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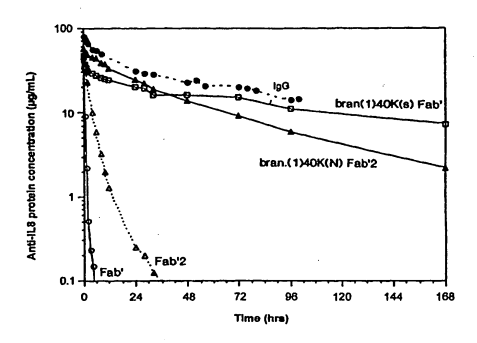
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(57) Abstract

Humanized anti-IL-8 monoclonal antibodies and variants thereof are described for use in diagnostic applications and in the treatment of inflammatory disorders. Also described is a conjugate formed by an antibody fragment covalently attached to a non-proteinaceous polymer, wherein the apparent size of the conjugate is at least about 500 kD. The conjugate exhibits substantially improved half-life, mean residence time, and/or clearance rate in circulation as compared to the underivatized parental antibody fragment.

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# ANTIBODY FRAGMENT-POLYMER CONJUGATES AND HUMANIZED ANTI-IL-8 MONOCLONAL ANTIBODIES

#### FIELD OF THE INVENTION

This application relates to the field of antibody fragments derivatized with polymers, and in particular to the use of such derivatization to increase the circulation half-lives of antibody fragment-polymer conjugates. This application also relates to humanized anti-interleukin-8 (IL-8) antibodies and to high affinity variants of such antibodies.

#### **BACKGROUND**

Modification of proteins with polyethylene glycol ("PEGylation") has the potential to increase residence time and reduce immunogenicity in vivo. For example, Knauf et al., J. Biol. Chem., 263: 15064-15070 (1988) reported a study of the pharmacodynamic behavior in rats of various polyoxylated glycerol and polyethylene glycol modified species of interleukin-2. Despite the known advantage of PEGylation, PEGylated proteins have not been widely exploited for clinical applications. In the case of antibody fragments, PEGylation has not been shown to extend serum half-life to useful levels. Delgado et al., Br. J. Cancer, 73: 175-182 (1996), Kitamura et al., Cancer Res., 51: 4310-4315 (1991), Kitamura et al., Biochem. Biophys. Res. Comm., 171: 1387-1394 (1990), and Pedley et al., Br. J. Cancer, 70: 1126-1130 (1994) reported studies characterizing blood clearance and tissue uptake of certain anti-tumor antigen antibodies or antibody fragments derivatized with low molecular weight (5 kD) PEG. Zapata et al., FASEB J., 9: A1479 (1995) reported that low molecular weight (5 or 10 kD) PEG attached to a sulfhydryl group in the hinge region of a Fab' fragment reduced clearance compared to the parental Fab' molecule.

Interleukin-8 (IL-8) is neutrophil chemotactic peptide secreted by a variety of cells in response to inflammatory mediators (for a review see Hebert et al. <u>Cancer Investigation</u> 11(6):743 (1993)). IL-8 can play an important role in the pathogenesis of inflammatory disorders, such as adult respiratory distress syndrome (ARDS), septic shock, and multiple organ failure. Immune therapy for such inflammatory disorders can include treatment of an affected patient with anti-IL-8 antibodies.

Sticherling et al. (J. Immunol. 143:1628 (1989)) disclose the production and characterization of four monoclonal antibodies against IL-8. WO 92/04372, published March 19, 1992, discloses polyclonal antibodies which react with the receptor-interacting site of IL-8 and peptide analogs of IL-8, along with the use of such antibodies to prevent an inflammatory response in patients. St. John et al. (Chest 103:932 (1993)) review immune therapy for ARDS, septic shock, and multiple organ failure, including the potential therapeutic use of anti-IL-8 antibodies. Sekido et al. (Nature 365:654 (1993)) disclose the prevention of lung reperfusion injury in rabbits by a monoclonal antibody against IL-8. Mulligan et al. (J. Immunol. 150:5585 (1993)), disclose protective effects of a murine monoclonal antibody to human IL-8 in inflammatory lung injury in rats.

WO 95/23865 (International Application No. PCT/US95/02589 published September 8, 1995) demonstrates that anti-IL-8 monoclonal antibodies can be used therapeutically in the treatment of other inflammatory disorders, such as bacterial pneumonias and inflammatory bowel disease.

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Anti-IL-8 antibodies are additionally useful as reagents for assaying IL-8. For example, Sticherling et al. (Arch. Dermatol. Res. 284:82 (1992)), disclose the use of anti-IL-8 monoclonal antibodies as reagents in immunohistochemical studies. Ko et al. (J. Immunol. Methods 149:227 (1992)) disclose the use of anti-IL-8 monoclonal antibodies as reagents in an enzyme-linked immunoabsorbent assay (ELISA) for IL-8.

### SUMMARY OF THE INVENTION

One aspect of the invention is a conjugate consisting essentially of one or more antibody fragments covalently attached to one or more polymer molecules, wherein the apparent size of the conjugate is at least about 500 kD.

Another aspect of the invention is an anti-IL-8 monoclonal antibody or antibody fragment comprising the complementarity determining regions of the 6G4.2.5LV11N35E light chain polypeptide amino acid sequence of Fig. 45 (SEQ ID NO: ).

Further aspects of the invention are a nucleic acid molecule comprising a nucleic acid sequence encoding the above-described anti-IL-8 monoclonal antibody or antibody fragment; an expression vector comprising the nucleic acid molecule operably linked to control sequences recognized by a host cell transfected with the vector; a host cell transfected with the vector; and a method of producing the antibody fragment comprising culturing the host cell under conditions wherein the nucleic acid encoding the antibody fragment is expressed, thereby producing the antibody fragment, and recovering the antibody fragment from the host cell.

#### BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a graph depicting the blocking of IL-8 mediated elastase release from neutrophils by anti-IL-8 monoclonal antibody 5.12.14.

Figure 2 is a graph depicting the inhibition of <sup>125</sup>l-IL-8 binding to neutrophils by unlabeled IL-8.

Figure 3 demonstrates that a isotype matched negative control Fab (denoted as "4D5 Fab") does not inhibit the binding of <sup>125</sup>I-IL-8 to human neutrophils.

Figure 4 is a graph depicting the inhibition of binding of  $^{125}$ I-IL-8 to human neutrophils by chimeric 5.12.14 Fab with an average IC<sub>50</sub> of 1.6 nM.

Figure 5 is a graph depicting the inhibition of binding of  $^{125}$ I-IL-8 to human neutrophils by chimeric 6G.4.25 Fab with an average IC<sub>50</sub> of 7.5 nM.

Figure 6 demonstrates the inhibition of human IL-8 mediated neutrophil chemotaxis by chimeric 6G4.2.5 Fab and chimeric 5.12.14 Fab.

Figure 7 demonstrates the relative abilities of chimeric 6G4.2.5 Fab and chimeric 5.12.14 Fab to inhibit rabbit IL-8 mediated neutrophil chemotaxis.

Figure 8 depicts the stimulation of elastase release from human neutrophils by various concentrations of human and rabbit IL-8. The relative extent of elastase release was quantitated by measurement of absorbance at 405 nm. The data represent mean ± SEM of triplicate samples.

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Figure 9 is a graph depicting the ability of chimeric 6G4.2.5 Fab and chimeric 5.12.14 Fab to inhibit elastase release from human neutrophils stimulated by human IL-8. The results were normalized to reflect the percentage of elastase release elicited by 100 nM IL-8 alone. The data represent the mean  $\pm$  SEM of three separate experiments performed on different days with different blood donors. IC<sub>50</sub> values were calculated by four parameter fit.

Figure 10 is a graph depicting the relative abilities of chimeric 6G4.2.5 Fab and chimeric 5.12.14 Fab to inhibit elastase release from human neutrophils stimulated by rabbit IL-8. The results were normalized to reflect the percentage of elastase release elicited by 100 nM IL-8 alone. The data represent the mean  $\pm$  SEM of three separate experiments performed on different days with different blood donors. IC<sub>50</sub> values were calculated by four parameter fit.

Figures 11A-11J are a set of graphs depicting the following parameters in a rabbit ulcerative colitis model: Figure 11A depicts myeloperoxidase levels in tissue; Figure 11B depicts IL-8 levels in tissue; Figure 11C depicts colon weight; Figure 11D depicts gross inflammation; Figure 11E depicts edema; Figure 11F depicts extent of necrosis; Figure 11G depicts severity of necrosis; Figure 11H depicts neutrophil margination; Figure 11I depicts neutrophil infiltration; and Figure 11J depicts mononuclear infiltration.

Figure 12 is a graph depicting the effect of anti-IL-8 monoclonal antibody treatment on the number of neutrophils in bronchoalveolar lavage (BAL) fluid in animals infected with <u>Streptococcus pneumoniae</u>, <u>Escherichia coli</u>, or <u>Pseudomonas aeruginosa</u>. Treatment with 6G4.2.5 significantly reduced the number of neutrophils present in the BAL fluid compared to animals treated with isotype control mouse IgG (Figure 12).

Figure 13 depicts the DNA sequences (SEQ ID NOS: 1-6) of three primers designed for each of the light and heavy chains. Multiple primers were designed in order to increase the chances of primer hybridization and efficiency of first strand cDNA synthesis for cloning the variable light and heavy regions of monoclonal antibody 5.12.14.

Figure 14 depicts the DNA sequences (SEQ ID NOS: 7-10) of one forward primer and one reverse primer for the 5.12.14 light chain variable region amplification.

Figure 15 depicts the DNA sequences (SEQ ID NOS: 11-18) of one forward primer and one reverse primer for the 5.12.14 heavy chain variable region amplification.

Figure 16 depicts the DNA sequence (SEQ ID NO: 19) and the amino acid sequence (SEQ ID NO: 20) of the 5.12.14 light chain variable region and partial murine constant light region. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). Important restriction sites are indicated in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable light region is amino acids 1 to 109. The partial murine constant light region is amino acids 110 to 123 (in italics).

Figure 17 depicts the DNA sequence (SEQ ID NO: 21) and the amino acid sequence (SEQ ID NO: 22) of the 5.12.14 heavy chain variable region and partial murine constant heavy region. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison

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(amino acids denoted with asterisk). Important restriction sites are indicated in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable heavy region is amino acids 1 to 120. The partial murine constant heavy region is amino acids 121 to 130.

Figure 18 depicts the DNA sequences (SEQ ID NOS: 23-26) of amplification primers used to convert murine light and heavy chain constant region residues to their human equivalents.

Figure 19 depicts the DNA sequence (SEQ ID NO: 27) and the amino acid sequence (SEQ ID NO: 28) for the 5.12.14 light chain variable region and the human IgG1 light chain constant region. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). The human constant region is denoted in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable light region is amino acids 1 to 109. The human constant light region is amino acids 110 to 215.

Figures 20A-20B depict the DNA sequence (SEQ ID NO: 29) and the amino acid sequence (SEQ ID NO: 30) for the 5.12.14 heavy chain variable region and the heavy chain constant region of human IgG1. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). The human constant region is denoted in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable heavy region is amino acids 1 to 120. The human constant heavy region is amino acids 121 to 229.

Figure 21 depicts the DNA sequences (SEQ ID NOS: 31-36) of three primers designed for each of the light and heavy chains. Multiple primers were designed in order to increase the chances of primer hybridization and efficiency of first strand cDNA synthesis for cloning the variable light and heavy regions of monoclonal antibody 6G4.2.5.

Figure 22 depicts the DNA sequences (SEQ ID NOS: 37-40) of one forward primer and one reverse primer for the 6G4.2.5 light chain variable region amplification.

Figure 23 depicts the DNA sequences (SEQ ID NOS: 41-46) of one forward primer and one reverse primer for the 6G4.2.5 heavy chain variable region amplification.

Figure 24 depicts the DNA sequence (SEQ ID NO: 47) and the amino acid sequence (SEQ ID NO: 48) of the 6G4.2.5 light chain variable region and partial murine constant light region. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). Useful cloning sites are in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable light region is amino acids 1 to 114. The partial murine constant light region is amino acids 115 to 131.

Figure 25 depicts the DNA sequence (SEQ ID NO: 49) and the amino acid sequence (SEQ ID NO: 50) of the 6G4.2.5 heavy chain variable region and partial murine constant heavy region. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). Useful cloning sites are in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable heavy region is amino acids 1 to 122. The partial murine constant heavy region is amino acids 123 to 135.

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Figure 26 depicts the DNA sequences (SEQ ID NOS: 51-54) of primers to convert the murine light chain and heavy chain constant regions to their human equivalents.

Figures 27A-27B depict the DNA sequence (SEQ ID NO: 55) and the amino acid sequence (SEQ ID NO: 56) for the chimeric 6G4.2.5 light chain. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). The human constant region is denoted in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable heavy region is amino acids 1 to 114. The human constant heavy region is amino acids 115 to 220.

Figures 28A-28B depict the DNA sequence (SEQ ID NO: 57) and the amino acid sequence (SEQ ID NO: 58) for the chimeric 6G4.2.5 heavy chain. CDRs are indicated by either X-ray crystallography (underlined amino acids) or by Kabat sequence comparison (amino acids denoted with asterisk). The human constant region is denoted in italics. The signal peptide of STII is amino acids -23 to -1. The murine variable heavy region is amino acids 1 to 122. The human constant heavy region is amino acids 123 to 231.

Fig. 29 depicts an amino acid sequence alignment of murine 6G425 light chain variable domain (SEQ ID NO: 59), humanized 6G425 F(ab)-1 light chain variable domain (SEQ ID NO: 60), and human light chain xI consensus framework (SEQ ID NO: 61) amino acid sequences, and an amino acid sequence alignment of murine 6G425 heavy chain variable domain (SEQ ID NO: 62), humanized 6G425 F(ab)-1 heavy chain variable domain (SEQ ID NO: 63), and human IgG1 subgroup III heavy chain variable domain (SEQ ID NO: 64) amino acid sequences, used in the humanization of 6G425. Light chain CDRs are labeled L1, L2, L3; heavy chain CDRs are labeled H1, H2, and H3. = and + indicate CDR sequences as defined by X-ray crystallographic contacts and sequence hypervariability, respectively. # indicates a difference between the aligned sequences. Residue numbering is according to Kabat et al. Lower case lettering denotes the insertion of an amino acid residue relative to the humIII consensus sequence numbering.

Fig. 30 is a graph with three panels (A, B and C) depicting the ability of F(ab)-9 (humanized 6G4V11 Fab) to inhibit human wild type IL-8, human monomeric IL-8, and rhesus IL-8 mediated neutrophil chemotaxis, respectively. Panel A presents inhibition data for F(ab)-9 samples at concentrations of 0.06 nM, 6.25 nM, 12.5 nM, 25 nM, 50 nM, and 100 nM, for an isotype control antibody (denoted "4D5") sample at a concentration of 100 nM, and for a no antibody control sample, in the presence of 2nM human wild type IL-8. Panel B presents inhibition data for F(ab)-9 samples at concentrations of 6.25 nM, 12.5 nM, 25 nM, and 50 nM, for an isotype control antibody (denoted "4D5") sample at a concentration of 100 nM, and for a no antibody control sample, in the presence of 4 nM human monomeric IL-8 (denoted as "BD59" and as "monomeric IL-8"). Panel C presents inhibition data for F(ab)-9 samples at concentrations of 1 nM, 12.5 nM, 25 nM, and 50 nM, for an isotype control antibody (denoted "4D5") sample at a concentration of 100 nM, and for a no antibody control sample, in the presence of 2 nM rhesus IL-8. In addition, all panels A, B an C each presents data for a no IL-8 buffer control sample (denoted as "Buffer") in the respective inhibition assay.

Fig. 31A depicts the amino acid sequences of the humanized anti-IL-8 6G4.2.5V11 light chain in an N-terminal fusion with the STII leader peptide (SEQ ID NO: 65), the humanized anti-IL-8 6G4.2.5V11

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heavy chain in an N-terminal fusion with the STII leader peptide (SEQ ID NO: 66), and a peptide linker in a C-terminal fusion with M13 phage gene-III coat protein (SEQ ID NO: 67).

Fig. 31B depicts the nucleic acid sequence (SEQ ID NO: 68) and the translated amino acid sequence (SEQ ID NO: 65) of the humanized anti-IL-8 6G4.2.5V11 light chain in an N-terminal fusion with the STII leader peptide.

Fig. 31C depicts the amino acid sequences of the humanized anti-IL-8 6G4.2.5V19 light chain in an N-terminal fusion with the STII leader peptide (SEQ ID NO: 69), and the humanized anti-IL-8 6G4.2.5V19 heavy chain in an N-terminal fusion with the STII leader peptide (SEQ ID NO: 70).

Fig. 32 is a three dimensional computer model of the humanized anti-IL-8 6G4.2.5V11 antibody. Heavy chain CDR loops and variable domain regions appear in purple, and CDR-H3 side chain residues appear in yellow. Heavy chain constant domain regions appear in red. Light chain CDR loops and variable domain regions appear in off-white, and the Asn residue at amino acid position 35 (N35) in CDR L1 appears in green. Light chain constant domain regions appear in amber.

Fig. 33 is a Scatchard plot depicting the inhibition of <sup>125</sup>I-IL-8 binding to human neutrophils exhibited by intact murine 6G4.2.5 antibody (denoted 6G4 murine mAb), 6G4.2.5 murine-human chimera Fab (denoted 6G4 chimera), humanized 6G4.2.5 Fab versions 1 and 11 (denoted V1 and V11), and variant 6G4.2.5V11N35A Fab (denoted V11N35A).

Fig. 34 is a graph with four panels (A, B, C, and D) depicting the ability of 6G4.2.5V11N35A Fab to inhibit human wild type IL-8, human monomeric IL-8, rabbit IL-8, and rhesus IL-8 mediated neutrophil Panel A presents inhibition data for 6G4.2.5V11N35A Fab samples at chemotaxis, respectively. concentrations of 0.5, 1, 2, 4, 8, 16, and 33 nM, for an isotype control antibody (denoted "4D5") sample at a concentration of 33 nM, and for a no antibody control (denoted "Hull-8") sample, in the presence of 2 nM human wild type IL-8. Panel B presents inhibition data for 6G4.2.5V11N35A Fab samples at concentrations of 0.5, 1, 2, 4, 8, 16, and 33 nM, for an intact 6G4.2.5 mAb sample at a concentration of 33 nM, for an isotype control antibody (denoted as "4D5") sample at a concentration of 33 nM, and for a no antibody control (denoted "BD59") sample, in the presence of 2 nM human monomeric IL-8. Panel C presents inhibition data for 6G4.2.5V11N35A Fab samples at concentrations of 0.5, 1, 2, 4, 8, 16, and 33 nM, for an intact 6G4.2.5 mAb sample at a concentration of 33 nM, for an isotype control antibody (denoted "4D5") sample at a concentration of 33 nM, and for a no antibody control (denoted "Rab IL-8") sample, in the presence of 2 nM rabbit 1L-8. Panel D presents inhibition data for 6G4.2.5V11N35A Fab samples at concentrations of 0.5, 1, 2, 4, 8, 16, and 33 nM, for an intact 6G4.2.5 mAb sample at a concentration of 33 nM, for an isotype control antibody (denoted as "4D5") sample at a concentration of 33 nM, and for a no antibody control (denoted "Rhe IL-8") sample, in the presence of 2 nM rhesus IL-8. In addition, panels B, C and D each presents data for human wild type IL-8 control (denoted "HulL-8") samples at a concentration of 2 nM in the respective assay, and panels A, B, C, and D each presents data for a no IL-8 buffer control (denoted "Buffer") sample in the respective assay.

Fig. 35 depicts the amino acid sequences of the humanized anti-IL-8 6G4.2.5V11N35A light chain

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in an N-terminal fusion with the STII leader peptide (SEQ ID NO: 71), the humanized anti-IL-8 6G4.2.5V11N35A heavy chain in an N-terminal fusion with the STII leader peptide (SEQ ID NO: 66), and the GCN4 leucine zipper peptide (SEQ ID NO: 72). The Ala residue (substituted for the wild type Asn residue) at amino acid position 35 in the 6G4.2.5V11N35A light chain appears in bold case. A putative pepsin cleavage site in the GCN4 leucine zipper sequence is underlined.

Fig. 36 depicts the DNA sequence (SEQ ID NO: 73) and the amino acid sequence (SEQ ID NO: 71) of the humanized anti-IL-8 6G4.2.5V11N35A light chain in an N-terminal fusion with the STII leader peptide. Complementarity determining regions L1, L2, and L3 are underlined

Figs. 37A-37B depict the DNA sequence (SEQ ID NO: 74) and the amino acid sequence (SEQ ID NO: 75) of the humanized anti-IL-8 6G4.2.5V11N35A heavy chain in an N-terminal fusion with the STII leader peptide and in a C-terminal fusion with the GCN4 leucine zipper sequence. Complementarity determining regions H1, H2, and H3 are underlined.

Fig. 38 is a Scatchard plot depicting the inhibition of <sup>125</sup>I-IL-8 binding to human neutrophils exhibited by 6G4.2.5V11N35A Fab (denoted Fab), 6G4.2.5V11N35A F(ab')<sub>2</sub> (denoted F(ab')<sub>2</sub>), and human wild type IL-8 control (denoted IL-8).

Fig. 39 is a graph depicting a comparison of the wild type human IL-8 mediated neutrophil chemotaxis inhibition activities of the 6G4.2.5V11N35A F(ab')<sub>2</sub> and 6G4.2.5V11N35A Fab. Inhibition data are presented for 6G4.2.5V11N35A Fab samples (denoted "N35A Fab") and 6G4.2.5V11N35A F(ab')<sub>2</sub> samples (denoted N35A F(ab')<sub>2</sub>) at concentrations of 0.3, 1, 3, 10, 30, and 100 nM, for an isotype control antibody (denoted as "4D5") sample at a concentration of 100 nM, and for a no antibody control sample, in the presence of 2 nM human wild type 1L-8. In addition, inhibition data are presented for no IL-8 buffer control samples (denoted "Buffer").

Fig. 40 is a graph depicting the ability of 6G4.2.5V11N35A F(ab')<sub>2</sub> to inhibit human monomeric IL-8, rhesus IL-8, and rabbit IL-8 mediated neutrophil chemotaxis. Human monomeric IL-8 mediated neutrophil chemotaxis data are presented for 6G4.2.5V11N35A F(ab')<sub>2</sub> samples at concentrations of 0.3, 1, 3, and 10 nM, for an isotype control antibody (denoted as "4D5") sample at a concentration of 100 nM, and for a no antibody control sample (denoted as "BD59"), in the presence of human monomeric IL-8 (denoted as "BD59") at a concentration of 0.5 nM. Rhesus IL-8 mediated neutrophil chemotaxis data are presented for 6G4.2.5V11N35A F(ab')<sub>2</sub> samples at concentrations of 0.3, 1, 3, and 10 nM, and for a no antibody control sample, in the presence of rhesus IL-8 at a concentration of 2 nM. Rabbit IL-8 mediated neutrophil chemotaxis data are presented for 6G4.2.5V11N35A F(ab')<sub>2</sub> samples at concentrations of 0.3, 1, 3, and 10 nM, and for a no antibody control sample, in the presence of rabbit IL-8 at a concentration of 2 nM. In addition, inhibition data are presented for a no IL-8 buffer control sample (denoted as "Buffer") and for a 2 nM human wild type IL-8 (denoted as "HuIL-8").

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Figs. 41A-41Q depict the nucleic acid sequence (SEQ ID NO: 76) of the p6G4V11N35A.F(ab')<sub>2</sub> vector.

Fig. 42 depicts the nucleic acid sequences of the stop template primer (SEQ ID NO: ) and the NNS randomization primer (SEQ ID NO: ) used for random mutagenesis of amino acid position 35 in variable light chain CDR-L1 of humanized antibody 6G4V11.

Fig. 43A is a table of data describing the frequencies of different phage display clones obtained from the randomization of amino acid position 35 in variable light chain CDR-L1 of humanized antibody 6G4V11.

Fig. 43B contains graphs of displacement curves depicting the inhibition of <sup>125</sup>I-IL-8 binding to neutrophils exhibited by the 6G4V11N35A, 6G4V11N35D, 6G4V11N35E and 6G4V11N35G Fab's.

Fig. 44 contains a graph depicting the typical kinetics of an anti-IL-8 antibody fragment (6G4V11N35A F(ab')2) binding to IL-8. Fig. 44 also contains a table of data providing the equilibrium constant for 6G4V11N35A Fab binding to IL-8 (rate constants were not determined "ND"), and the equilibrium and rate constants for 6G4V11N35A F(ab')2 and 6G4V11N35E Fab binding to IL-8.

Fig. 45 depicts the DNA sequence (SEQ ID NO: ) and amino acid sequence (SEQ ID NO: ) of the 6G4V11N35E light chain in an N-terminal fusion with the STII leader peptide. Complementarity determining regions L1, L2 and L3 are underlined.

Fig. 46 is a graph depicting the ability of 6G4V11N35E Fab to inhibit human IL-8 (dark columns) and rabbit IL-8 (light columns) mediated neutrophil chemotaxis. Data are presented for 6G4V11N35E Fab samples at concentrations of 0.4, 1.2, 3.7, 11 and 33 nM, and for an isotype control antibody (4D5) sample at a concentration of 100 nM, in the presence of 2 nM human IL-8 or 2 nM rabbit IL-8. In addition, inhibition data are presented for a no IL-8 buffer control sample (denoted "Buffer") and for human and rabbit IL-8 control samples (denoted "IL-8").

Fig. 47 depicts the DNA sequence of the sense (SEQ ID NO: ) and anti-sense (SEQ ID NO: ) strands of a PvuII-XhoI synthetic nucleotide encoding amino acids Leu4 to Phe29 of the 6G4V11N35A heavy chain.

Figs. 48A-48T depict the DNA sequence (SEQ ID NO: ) of plasmid p6G4V11N35A.choSD9.

Fig. 49 contains graphs of displacement curves depicting the inhibition of <sup>125</sup>I-IL-8 binding to neutrophils exhibited by the full length IgG1 forms of variants 6G4V11N35A and 6G4V11N35E.

Figs. 50A-50B are graphs depicting the ability of full length 6G4V11N35A IgG1 and 6G4V11N35E IgG1 to inhibit human IL-8 (Fig. 50A) and rabbit IL-8 (Fig. 50B) mediated neutrophil chemotaxis.

Fig. 51 contains a graph depicting the typical kinetics of a full length anti-IL8 antibody (6G4V11N35A IgG1) binding to IL-8. Fig. 51 also contains a table of data providing the equilibrium and rate constants for full length murine 6G4.2.5 IgG2a, 6G4V11N35A IgG1 and 6G4V11N35E IgG1 binding to IL-8.

Fig. 52 contains graphs of displacement curves depicting the results of an unlabeled IL-8/<sup>125</sup>I-IL-8 competition radioimmunoassay performed with full length 6G4V11N35A IgG1 and 6G4V11N35E IgG1.

Fig. 53 depicts the DNA sequence (SEQ ID NO: ) and amino acid sequence (SEQ ID NO: ) of the 6G4V11N35A Fab' heavy chain (6G4V11N35A Fab heavy chain modified to contain a cysteine residue in the hinge region).

Figs. 54A-54C contain graphs of displacement curves depicting the IL-8 binding and IC<sub>50</sub>'s for PEG-maleimide modified 6G4V11N35A Fab' molecules.

Figs. 55A-55C are graphs depicting the ability of PEG-maleimide modified 6G4V11N35A Fab' molecules to inhibit human IL-8 and rabbit IL-8 mediated neutrophil chemotaxis.

Figs. 56A-56C are graphs depicting the ability of PEG-maleimide modified 6G4V11N35A Fab' molecules to inhibit IL-8 mediated release of  $\beta$ -glucuronidase from neutrophils.

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Figs. 57A-57B contain graphs of displacement curves depicting the inhibition of <sup>125</sup>I-IL-8 binding to neutrophils exhibited by PEG-succinimide modified 6G4V11N35A Fab'<sub>2</sub> molecules.

Figs. 58A-58B are graphs depicting the ability of PEG-succinimide modified 6G4V11N35A F(ab')<sub>2</sub> molecules to inhibit human IL-8 mediated neutrophil chemotaxis.

Figs. 59A-59B are graphs depicting the ability of PEG-succinimide modified 6G4V11N35A  $F(ab')_2$  molecules to inhibit human IL-8 mediated release of  $\beta$ -glucuronidase from neutrophils.

Fig. 60 is a graph depicting the theoretical molecular weight (dotted bars) and effective size (solid bars) of PEG-maleimide modified 6G4V11N35A Fab' molecules as determined by SEC-HPLC.

Fig. 61 is an SDS-PAGE gel depicting the electrophoretic mobility of various PEG-maleimide modified 6G4V11N35A Fab' molecules.

Fig. 62 contains size exclusion chromatograms (SEC-HPLC) depicting the retention times and effective (hydrodynamic) sizes of various PEG-succinimide modified 6G4V11N35A F(ab')<sub>2</sub> molecules.

Fig. 63 is a graph depicting the theoretical molecular weight (open columns), effective size determined by SEC-HPLC (solid columns), and the actual molecular weight determined by SEC-light scattering (shaded columns) for various PEG-succinimide modified 6G4V11N35A F(ab')<sub>2</sub> molecules.

Fig. 64 is an SDS-PAGE gel depicting the electrophoretic mobility of various PEG-succinimide modified 6G4V11N35A F(ab')<sub>2</sub> molecules. From left to right, lane 1 contains unmodified F(ab')<sub>2</sub>, lane 2 contains F(ab')<sub>2</sub> coupled to two 40 kD branched PEG-succinimide molecules (denoted "Br(2)-40kD(N)-F(ab')2"), lane 3 contains F(ab')<sub>2</sub> coupled to one 40 kD branched PEG-succinimide molecule (denoted "Br(1)-40kD-(N)-Fab'2"), lane 4 contains a mixture of F(ab')<sub>2</sub> coupled to four 20 kD linear PEG-succinimide molecules and F(ab')<sub>2</sub> c upled to five 20 kD linear PEG-succinimide molecules (denoted

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"L(4+5)-20kD-(N)-Fab'2"), lane 5 contains F(ab')<sub>2</sub> coupled to one 20 kD linear PEG-succinimide molecule (denoted "L(1)-20kD-(N)-Fab'2"), and lane 6 contains molecular weight standards.

Fig. 65 contains graphs comparing the serum concentration vs. time profiles of various PEG-maleimide modified 6G4V11N35A Fab' molecules (upper graph) and various PEG-succinimide modified 6G4V11N35A F(ab')<sub>2</sub> molecules (lower graph) in rabbits. In the upper graph, "bran.(1)40K(s)Fab' "denotes 6G4V11N35A Fab' coupled to one 40 kD branched PEG-maleimide molecule, "lin.(1)40K(s)Fab' "denotes 6G4V11N35A Fab' coupled to one 40 kD linear PEG-maleimide molecule, "lin.(1)30K(s)Fab' "denotes 6G4V11N35A Fab' coupled to one 30 kD linear PEG-maleimide molecule, "lin.(1)20K(s)Fab'' denotes 6G4V11N35A Fab' coupled to one 20 kD linear PEG-maleimide molecule. In the lower graph, "bran.(2)40K(N)Fab'2" denotes 6G4V11N35A F(ab')<sub>2</sub> coupled to two 40 kD branched PEG-succinimide molecules, "bran.(1)40K(N)Fab'2" denotes 6G4V11N35A F(ab')<sub>2</sub> coupled to one 40 kD branched PEG-succinimide molecule, and "Fab'2" denotes unmodified 6G4V11N35A F(ab')<sub>2</sub>. In both graphs, "lgG" denotes a full length lgG1 equivalent of the human-murine chimeric anti-rabbit IL-8 Fab described in Example F below.

Fig. 66 contains graphs comparing the serum concentration vs. time profiles of 6G4V11N35A Fab' coupled to one 40 kD branched PEG-maleimide molecule (denoted as "bran.(1)40K(s)Fab"), 6G4V11N35A F(ab')<sub>2</sub> coupled to one 40 kD branched PEG-succinimide molecule (denoted as "bran.(1)40K(N)Fab'2"), unmodified 6G4V11N35A F(ab')<sub>2</sub> (denoted as "Fab'2"), unmodified 6G4V11N35A Fab' (denoted as "Fab"), and a full length IgG1 (denoted as "IgG") equivalent of the human-murine chimeric anti-rabbit IL-8 Fab described in Example F below.

Fig. 67 is a graph depicting the effect of 6G4V11N35A Fab' coupled to one 40 kD branched PEG-maleimide molecule (denoted as "PEG 40 Kd") and murine anti-rabbit IL-8 monoclonal antibody 6G4.2.5 (full length IgG2a) (denoted as "6G4.2.5") on gross weight of entire lung in an ARDS rabbit model.

Fig. 68 is a graph depicting the effect of 6G4V11N35A Fab' coupled to one branched 40 kD PEG-maleimide molecule (denoted as "PEG 40 Kd") and murine anti-rabbit IL-8 monoclonal antibody 6G4.2.5 (full length IgG2a) (denoted as "6G4.2.5") on BAL total leukocyte (light columns) and polymorphonuclear cell (dark columns) counts in an ARDS rabbit model. Untreated (no therapeutics) control animal data is denoted as "Control".

Fig. 69 is a graph depicting the effect of 6G4V11N35A Fab' coupled to one branched 40 kD PEG-maleimide molecule (denoted as "PEG 40 Kd") and murine anti-rabbit 1L-8 monoclonal antibody 6G4.2.5 (full length IgG2a) (denoted as "6G4.2.5") on PaO2/FiO2 ratio at 24 hours-post treatment (light columns) and 48 hours post-treatment (dark columns) in an ARDS rabbit model. Untreated (no therapeutics) control animal data is denoted as "Control".

#### DESCRIPTION OF THE PREFERRED EMBODIMENTS

#### I. DEFINITIONS

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In general, the following words or phrases have the indicated definition when used in the description, examples, and claims.

"Polymerase chain reaction" or "PCR" refers to a procedure or technique in which minute amounts of a specific piece of nucleic acid, RNA and/or DNA, are amplified as described in U.S. Patent No. 4,683,195 issued 28 July 1987. Generally, sequence information from the ends of the region of interest or beyond needs to be available, such that oligonucleotide primers can be designed; these primers will be identical or similar in sequence to opposite strands of the template to be amplified. The 5' terminal nucleotides of the two primers can coincide with the ends of the amplified material. PCR can be used to amplify specific RNA sequences, specific DNA sequences from total genomic DNA, and cDNA transcribed from total cellular RNA, bacteriophage or plasmid sequences, etc. See generally Mullis et al., Cold Spring Harbor Symp. Quant. Biol. 51:263 (1987); Erlich, ed., PCR Technology (Stockton Press, NY, 1989). As used herein, PCR is considered to be one, but not the only, example of a nucleic acid polymerase reaction method for amplifying a nucleic acid test sample comprising the use of a known nucleic acid as a primer and a nucleic acid polymerase to amplify or generate a specific piece of nucleic acid.

"Antibodies" (Abs) and "immunoglobulins" (Igs) are glycoproteins having the same structural characteristics. While antibodies exhibit binding specificity to a specific antigen, immunoglobulins include both antibodies and other antibody-like molecules which lack antigen specificity. Polypeptides of the latter kind are, for example, produced at low levels by the lymph system and at increased levels by myelomas.

"Native antibodies and immunoglobulins" are usually heterotetrameric glycoproteins of about 150,000 daltons, composed of two identical light (L) chains and two identical heavy (H) chains. Each light chain is linked to a heavy chain by one covalent disulfide bond, while the number of disulfide linkages varies between the heavy chains of different immunoglobulin isotypes. Each heavy and light chain also has regularly spaced intrachain disulfide bridges. Each heavy chain has at one end a variable domain (V<sub>H</sub>) followed by a number of constant domains. Each light chain has a variable domain at one end (V<sub>L</sub>) and a constant domain at its other end; the constant domain of the light chain is aligned with the first constant domain of the heavy chain, and the light chain variable domain is aligned with the variable domain of the heavy chain. Particular amino acid residues are believed to form an interface between the light- and heavy-chain variable domains (Clothia et al., J. Mol. Biol. 186:651 (1985); Novotny and Haber, Proc. Natl. Acad. Sci. U.S.A. 82:4592 (1985)).

The term "variable" refers to the fact that certain portions of the variable domains differ extensively in sequence among antibodies and are used in the binding and specificity of each particular antibody for its particular antigen. However, the variability is not evenly distributed throughout the variable domains of antibodies. It is concentrated in three segments called complementarity-determining regions (CDRs) or hypervariable regions both in the light-chain and the heavy-chain variable domains. The more highly

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conserved portions of variable domains are called the framework (FR). The variable domains of native heavy and light chains each comprise four FR regions, largely adopting a β-sheet configuration, connected by three CDRs, which form loops connecting, and in some cases forming part of, the β-sheet structure. The CDRs in each chain are held together in close proximity by the FR regions and, with the CDRs from the other chain, contribute to the formation of the antigen-binding site of antibodies (see Kabat *et al.*, Sequences of Proteins of Immunological Interest, Fifth Edition, National Institute of Health, Bethesda, MD (1991)). The constant domains are not involved directly in binding an antibody to an antigen, but exhibit various effector functions, such as participation of the antibody in antibody-dependent cellular toxicity.

Papain digestion of antibodies produces two identical antigen-binding fragments, called "Fab" fragments, each with a single antigen-binding site, and a residual "Fc" fragment, whose name reflects its ability to crystallize readily. Pepsin treatment yields an F(ab')<sub>2</sub> fragment that has two antigen-combining sites and is still capable of cross-linking antigen.

"Fv" is the minimum antibody fragment which contains a complete antigen-recognition and binding site. In a two-chain Fv species, this region consists of a dimer of one heavy- and one light-chain variable domain in tight, non-covalent association. In a single-chain Fv species (scFv), one heavy- and one light-chain variable domain can be covalently linked by a flexible peptide linker such that the light and heavy chains can associate in a "dimeric" structure analogous to that in a two-chain Fv species It is in this configuration that the three CDRs of each variable domain interact to define an antigen-binding site on the surface of the VH-VL dimer. Collectively, the six CDRs confer antigen-binding specificity to the antibody. However, even a single variable domain (or half of an Fv comprising only three CDRs specific for an antigen) has the ability to recognize and bind antigen, although at a lower affinity than the entire binding site. For a review of scFv see Pluckthun, in *The Pharmacology of Monoclonal Antibodies*, vol. 113, Rosenburg and Moore eds., Springer-Verlag, New York, pp. 269-315 (1994).

The Fab fragment also contains the constant domain of the light chain and the first constant domain (CH1) of the heavy chain. Fab' fragments differ from Fab fragments by the addition of a few residues at the carboxy terminus of the heavy chain CH1 domain including one or more cysteines from the antibody hinge region. Fab'-SH is the designation herein for Fab' in which the cysteine residue(s) of the constant domains bear a free thiol group. F(ab')<sub>2</sub> antibody fragments originally were produced as pairs of Fab' fragments which have hinge cysteines between them. Other chemical couplings of antibody fragments are also known.

The "light chains" of antibodies (immunoglobulins) from any vertebrate species can be assigned to one of two clearly distinct types, called kappa (k) and lambda (l), based on the amino acid sequences of their constant domains.

Depending on the amino acid sequence of the constant domain of their heavy chains, immunoglobulins can be assigned to different classes. There are five major classes of immunoglobulins: IgA, IgD, IgE, IgG, and IgM, and several of these can be further divided into subclasses (isotypes), e.g., IgG<sub>1</sub>, IgG<sub>2</sub>, IgG<sub>3</sub>, IgG<sub>4</sub>, IgA<sub>1</sub>, and IgA<sub>2</sub>. The heavy-chain constant domains that correspond to the different

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classes of immunoglobulins ar called  $\alpha$ ,  $\delta$ ,  $\epsilon$ ,  $\gamma$ , and  $\mu$ , respectively. The subunit structures and three-dimensional configurations of different classes of immunoglobulins are well known.

The term "antibody" is used in the broadest sense and specifically covers single monoclonal antibodies (including agonist and antagonist antibodies) and antibody compositions with polyepitopic specificity.

"Antibody fragment", and all grammatical variants thereof, as used herein are defined as a portion of an intact antibody comprising the antigen binding site or variable region of the intact antibody, wherein the portion is free of the constant heavy chain domains (i.e. CH2, CH3, and CH4, depending on antibody isotype) of the Fc region of the intact antibody. Examples of antibody fragments include Fab, Fab', Fab'-SH, F(ab')2, and Fv fragments; diabodies; any antibody fragment that is a polypeptide having a primary structure consisting of one uninterrupted sequence of contiguous amino acid residues (referred to herein as a "single-chain antibody fragment" or "single chain polypeptide"), including without limitation (1)single-chain Fv (scFv) molecules (2)single chain polypeptides containing only one light chain variable domain, or a fragment thereof that contains the three CDRs of the light chain variable domain, without an associated heavy chain moiety and (3)single chain polypeptides containing only one heavy chain variable region, or a fragment thereof containing the three CDRs of the heavy chain variable region, without an associated light chain moiety; and multispecific or multivalent structures formed from antibody fragments. In an antibody fragment comprising one or more heavy chains, the heavy chain(s) can contain any constant domain sequence (e.g. CH1 in the IgG isotype) found in a non-Fc region of an intact antibody, and/or can contain any hinge region sequence found in an intact antibody, and/or can contain a leucine zipper sequence fused to or situated in the hinge region sequence or the constant domain sequence of the heavy chain(s). Suitable leucine zipper sequences include the jun and fos leucine zippers taught by Kostelney et al., J. Immunol., 148: 1547-1553 (1992) and the GCN4 leucine zipper described in the Examples below.

Unless specifically indicated to the contrary, the term "conjugate" as described and claimed herein is defined as a heterogeneous molecule formed by the covalent attachment of one or more antibody fragment(s) to one or more polymer molecule(s), wherein the heterogeneous molecule is water soluble, i.e. soluble in physiological fluids such as blood, and wherein the heterogeneous molecule is free of any structured aggregate. In the context of the foregoing definition, the term "structured aggregate" refers to (1) any aggregate of molecules in aqueous solution having a spheroid or spheroid shell structure, such that the heterogeneous molecule is not in a micelle or other emulsion structure, and is not anchored to a lipid bilayer, vesicle or liposome; and (2) any aggregate of molecules in solid or insolubilized form, such as a chromatography bead matrix, that does not release the heterogeneous molecule into solution upon contact with an aqueous phase. Accordingly, the term "conjugate" as defined herein encompasses the aforementioned heterogeneous molecule in a precipitate, sediment, bioerodible matrix or other solid capable of releasing the heterogeneous molecule into aqueous solution upon hydration of the solid.

Unless specifically indicated to the contrary, the terms "polymer", "polymer molecule", "nonproteinaceous polymer", and "nonproteinaceous polymer molecule" are used interchangeably and are

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defined as a molecule formed by covalent linkage of two or more monomers, wherein none of the monomers is contained in the group consisting of alanine (Ala), cysteine (Cys), aspartic acid (Asp), glutamic acid (Glu), phenylalanine (Phe), glycine (Gly), histidine (His), isoleucine (Ile), lysine (Lys), leucine (Leu), methionine (Met), asparagine (Asn), proline (Pro), glutamine (Gln), arginine (Arg), serine (Ser), threonine (Thr), valine (Val), tryptophan (Trp), and tyrosine (Tyr) residues.

The term "monoclonal antibody" (mAb) as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally occurring mutations that may be present in minor Monoclonal antibodies are highly specific, being directed against a single antigenic site. Furthermore, in contrast to conventional (polyclonal) antibody preparations which typically include different antibodies directed against different determinants (epitopes), each mAb is directed against a single determinant on the antigen. In addition to their specificity, the monoclonal antibodies are advantageous in that they can be synthesized by hybridoma culture, uncontaminated by other immunoglobulins. The modifier "monoclonal" indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies, and is not to be construed as requiring production of the antibody by any particular method. For example, the monoclonal antibodies to be used in accordance with the present invention may be made by the hybridoma method first described by Kohler et al., Nature, 256:495 (1975), or may be made by recombinant DNA methods (see, e.g., U.S. Patent No. 4,816,567 to Cabilly et al.). The "monoclonal antibodies" also include clones of antigen-recognition and binding-site containing antibody fragments (Fv clones) isolated from phage antibody libraries using the techniques described in Clackson et al., Nature, 352:624-628 (1991) and Marks et al., J. Mol. Biol., 222:581-597 (1991), for example.

The monoclonal antibodies herein include hybrid and recombinant antibodies produced by splicing a variable (including hypervariable) domain of an anti-IL-8 antibody with a constant domain (e.g. "humanized" antibodies), or a light chain with a heavy chain, or a chain from one species with a chain from another species, or fusions with heterologous proteins, regardless of species of origin or immunoglobulin class or subclass designation, as well as antibody fragments (e.g., Fab, F(ab')<sub>2</sub>, and Fv), so long as they exhibit the desired biological activity. (See, e.g., U.S. Pat. No. 4,816,567 to Cabilly et al.; Mage and Lamoyi, in Monoclonal Antibody Production Techniques and Applications, pp. 79-97 (Marcel Dekker, Inc., New York, 1987).)

The monoclonal antibodies herein specifically include "chimeric" antibodies (immunoglobulins) in which a portion of the heavy and/or light chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a particular antibody class or subclass, while the remainder of the chain(s) is identical with or homologous to corresponding sequences in antibodies derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies, so long as they exhibit the desired biological activity (Cabilly et al., supra; Morrison et al., Proc. Natl. Acad. Sci. U.S.A. 81:6851 (1984)).

"Humanized" forms of non-human (e.g., murine) antibodies are specific chimeric

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immun globulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')<sub>2</sub>, or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. For the most part, humanized antibodies are human immunoglobulins (recipient antibody) in which residues from a complementary-determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat, or rabbit having the desired specificity, affinity, and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Furthermore, humanized antibodies can comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. These modifications are made to further refine and maximize antibody performance. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. For further details see Jones et al., Nature 321:522 (1986); Reichmann et al., Nature 332:323 (1988); and Presta. Curr. Op. Struct. Biol. 2:593 (1992).

"Treatment" refers to both therapeutic treatment and prophylactic or preventative measures. Those in need of treatment include those already with the disorder as well as those prone to have the disorder or those in which the disorder is to be prevented.

"Mammal" for purposes of treatment refers to any animal classified as a mammal, including humans, domestic and farm animals, and zoo, sports, or pet animals, such as dogs, horses, cats, cows, etc. Preferably, the mammal herein is human.

As used herein, protein, peptide and polypeptide are used interchangeably to denote an amino acid polymer or a set of two or more interacting or bound amino acid polymers.

As used herein, the term "inflammatory disorders" refers to pathological states resulting in inflammation, typically caused by neutrophil chemotaxis. Examples of such disorders include inflammatory skin diseases including psoriasis; responses associated with inflammatory bowel disease (such as Crohn's disease and ulcerative colitis); ischemic reperfusion: adult respiratory distress syndrome; dermatitis; meningitis; encephalitis; uveitis; autoimmune diseases such as rheumatoid arthritis, Sjorgen's syndrome, vasculitis; diseases involving leukocyte diapedesis; central nervous system (CNS) inflammatory disorder, multiple organ injury syndrome secondary to septicaemia or trauma; alcoholic hepatitis, bacterial pneumonia, antigen-antibody complex mediated diseases; inflammations of the lung, including pleurisy, alveolitis, vasculitis, pneumonia, chronic bronchitis, bronchiectasis, and cystic fibrosis; etc. The preferred indications are bacterial pneumonia and inflammatory bowel disease such as ulcerative colitis.

The terms "hydrodynamic size", "apparent size", "apparent molecular weight", "effective size" and "effective molecular weight" of a molecule are used synonymously herein refer to the size of a molecule as determined by comparison to a standard curve produced with globular protein molecular weight standards in a size exclusion chromatography system, wherein the standard curve is created by mapping the actual

molecular weight of each standard against its elution time bserved in the size exclusion chromatography system. Thus, the apparent size of a test molecule is derived by using the molecule's elution time to extrapolate a putative molecular weight from the standard curve. Preferably, the molecular weight standards used to create the standard curve are selected such that the apparent size of the test molecule falls within the linear portion of the standard curve.

#### II. MODES FOR CARRYING OUT THE INVENTION

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In one part, the invention arises from the surprising and unexpected discovery that antibody fragment-polymer conjugates having an effective or apparent size significantly greater than the antibody fragment-polymer conjugates described in the art confers an increase in serum half-life, an increase in mean residence time in circulation (MRT), and/or a decrease in serum clearance rate over underivatized antibody fragment which far exceed the modest changes in such biological property or properties obtained with the art-known antibody fragment-polymer conjugates. The present inventors have determined for the first time that increasing the effective size of an antibody fragment to at least about 500,000 D, or increasing the effective size of an antibody fragment by at least about 8 fold over the effective size of the parental antibody fragment, or derivatizing an antibody fragment with a polymer of at least about 20,000 D in molecular weight, yields a molecule with a commercially useful pharmacokinetic profile. The greatly extended serum half-life, extended MRT, and/or reduced serum clearance rate of the conjugates of the invention makes such conjugates viable alternatives to intact antibodies used for therapeutic treatment of many disease indications. Antibody fragments provide significant advantages over intact antibodies, notably the fact that recombinant antibody fragments can be made in bacterial cell expression systems. Bacterial cell expression systems provide several advantages over mammalian cell expression systems, including reduced time and cost at both the research and development and manufacturing stages of a product.

In another part, the present invention also arises from the humanization of the 6G4.2.5 murine antirabbit IL-8 monoclonal antibody ("6G4.2.5") described in WO 95/23865 (PCT/US95/02589 published September 8, 1995), the entire disclosure of which is specifically incorporated herein by reference. The hybridoma producing antibody 6G4.2.5 was deposited on September 28, 1994 with the American Type Culture Collection and assigned ATCC Accession No. HB 11722 as described in the Examples below. In one aspect, the invention provides a humanized derivative of the 6G4.2.5 antibody, variant 11 (referred to herein as "6G4.2.5v11"), in which the murine CDRs of 6G4.2.5 are grafted onto a consensus framework for human light chain x1 and human IgG1 heavy chain subgroup III, followed by importing three framework residues from the murine 6G4.2.5 parent heavy chain variable domain sequence into analogous sites in the heavy chain variable domain of the human template sequence, as described in the Examples below. In another aspect, the invention provides variants of the 6G4.2.5v11 antibody with certain amino acid substitution(s) yielding increased affinity for human IL-8 and/or promoting greater efficiency in recombinant manufacturing processes.

It will be understood that in the context of this Section (II) and all subsections thereof, every reference to "an antibody fragment" or "the antibody fragment" contained in a conjugate shall be a reference

to one or more antibody fragment(s) in the conjugate (consistent with the definition of the term "conjugate" set forth in Section (I) above), except where the number of antibody fragment(s) in the conjugate is expressly indicated. It will be understood that in the context of this Section (II) and all subsections thereof, every reference to "a polymer", "a polymer molecule", "the polymer", or "the polymer molecule" contained in a conjugate shall be a reference to one or more polymer molecule(s) in the conjugate (consistent with the definition of the term "conjugate" set forth in Section (I) above), except where the number of polymer molecule(s) in the conjugate is expressly indicated.

## 1. LARGE EFFECTIVE SIZE ANTIBODY FRAGMENT-POLYMER CONJUGATES

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In one aspect, the invention provides an antibody fragment covalently attached to a polymer to form a conjugate having an effective or apparent size of at least about 500,000 Daltons (D). In another aspect, the invention provides an antibody fragment covalently attached to a polymer to form a conjugate having an apparent size that is at least about 8 fold greater than the apparent size of the parental antibody fragment. In yet another aspect, the invention provides an antibody fragment covalently attached to a polymer of at least about 20,000 D in molecular weight (MW). It will be appreciated that the unexpectedly and surprisingly large increase in antibody fragment serum half-life, increase in MRT, and/or decrease in serum clearance rate can be achieved by using any type of polymer or number of polymer molecules which will provide the conjugate with an effective size of at least about 500,000 D, or by using any type of polymer or number of polymer molecules which will provide the conjugate with an effective size that is at least about 8 fold greater than the effective size of the parental antibody fragment, or by using any type or number of polymers wherein each polymer molecule is at least about 20,000 D in MW. Thus, the invention is not dependent on the use of any particular polymer or molar ratio of polymer to antibody fragment in the conjugate.

In addition, the beneficial aspects of the invention extend to antibody fragments without regard to antigen specificity. Although variations from antibody to antibody are to be expected, the antigen specificity of a given antibody will not substantially impair the extraordinary improvement in serum half-life, MRT, and/or serum clearance rate for antibody fragments thereof that can be obtained by derivatizing the antibody fragments as taught herein.

In one embodiment, the conjugate has an effective size of at least about 500,000 D, or at least about 800,000 D, or at least about 900,000 D, or at least about 1,200,000 D, or at least about 1,200,000 D, or at least about 1,400,000 D, or at least about 1,500,000 D, or at least about 2,000,000 D, or at least about 2,500,000 D.

In another embodiment, the conjugate has an effective size of at or about 500,000 D to at or about 10,000,000 D, or an effective size of at or about 500,000 D to at or about 8,000,000 D, or an effective size of at or about 500,000 D to at or about 500,000 D to at or about 4,000,000 D, or an effective size of at or about 3,000,000 D, or an effective size of at or about 500,000 D to at or about 2,000,000 D, or an effective size of at or about 2,000,000 D, or an effective size of at or about 1,800,000 D, or an effective size of at or about 1,800,000 D, or an

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effective size of at or about 500,000 D to at or about 1,600,000 D, or an effective size of at or about 500,000 D to at or about 1,500,000 D, or an effective size of at or about 500,000 D to at or about 1,000,000 D.

In another embodiment, the conjugate has an effective size of at or about 800,000 D to at or about 10,000,000 D, or an effective size of at or about 800,000 D to at or about 8,000,000 D, or an effective size of at or about 800,000 D to at or about 4,000,000 D, or an effective size of at or about 3,000,000 D, or an effective size of at or about 3,000,000 D, or an effective size of at or about 800,000 D to at or about 800,000 D to at or about 800,000 D to at or about 800,000 D, or an effective size of at or about 1,800,000 D, or an effective size of at or about 1,800,000 D, or an effective size of at or about 1,500,000 D, or an effective size of at or about 1,500,000 D.

In another embodiment, the conjugate has an effective size of at or about 900,000 D to at or about 10,000,000 D, or an effective size of at or about 900,000 D to at or about 8,000,000 D, or an effective size of at or about 900,000 D to at or about 900,000 D to at or about 4,000,000 D, or an effective size of at or about 900,000 D, or an effective size of at or about 900,000 D to at or about 2,500,000 D, or an effective size of at or about 900,000 D, or an effective size of at or about 1,800,000 D, or an effective size of at or about 1,500,000 D, or an effective size of at or about 1,500,000 D.

In another embodiment, the conjugate has an effective size of at or about 1,000,000 D to at or about 10,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D to at or about 1,000,000 D to at or about 1,000,000 D, or an effective size of at or about 2,500,000 D, or an effective size of at or about 1,000,000 D to at or about 1,000,000 D to at or about 1,000,000 D to at or about 1,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D, or an effective size of at or about 1,000,000 D to at or about 1,000,000 D to at or about 1,000,000 D to at or about 1,000,000 D.

In a further embodiment, the conjugate has an effective size that is at least about 8 fold greater, or at least about 10 fold greater, or at least about 12 fold greater, or at least about 15 fold greater, or at least about 18 fold greater, or at least about 20 fold greater, or at least about 25 fold greater, or at least about 28 fold greater, or at least about 30 fold greater, or at least about 40 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 8 fold to about 100 fold greater, or is about 8 fold to about 80 fold greater, or is about 8 fold to about 50 fold greater, or is about 8 fold to about 40 fold greater, or is about 8 fold to about 30 fold greater, or is about 8 fold to about 28 fold greater, or is about 8 fold to about 25 fold greater, or is about 8 fold to about 20 fold greater, or is about 8 fold to about 18 fold greater, or is about 8 fold to about 15 fold greater, than the effective size of the parental antibody fragment.

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In another embodiment, the conjugate has an effective size that is about 12 fold to about 100 fold greater, or is about 12 fold to about 80 fold greater, or is about 12 fold to about 50 fold greater, or is about 12 fold to about 40 fold greater, or is about 12 fold to about 30 fold greater, or is about 12 fold to about 28 fold greater, or is about 12 fold to about 25 fold greater, or is about 12 fold to about 20 fold greater, or is about 12 fold to about 18 fold greater, or is about 15 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 15 fold to about 100 fold greater, or is about 15 fold to about 80 fold greater, or is about 15 fold to about 50 fold greater, or is about 15 fold to about 40 fold greater, or is about 15 fold to about 30 fold greater, or is about 15 fold to about 28 fold greater, or is about 15 fold to about 25 fold greater, or is about 15 fold to about 20 fold greater, or is about 15 fold to about 18 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 18 fold to about 100 fold greater, or is about 18 fold to about 80 fold greater, or is about 18 fold to about 50 fold greater, or is about 18 fold to about 40 fold greater, or is about 18 fold to about 30 fold greater, or is about 18 fold to about 28 fold greater, or is about 18 fold to about 25 fold greater, or is about 18 fold to about 20 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 20 fold to about 100 fold greater, or is about 20 fold to about 80 fold greater, or is about 20 fold to about 50 fold greater, or is about 20 fold to about 40 fold greater, or is about 20 fold to about 30 fold greater, or is about 20 fold to about 28 fold greater, or is about 20 fold to about 25 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 25 fold to about 100 fold greater, or is about 25 fold to about 25 fold to about 50 fold greater, or is about 25 fold to about 40 fold greater, or is about 25 fold to about 30 fold greater, or is about 25 fold to about 25 fold to about 25 fold to about 25 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 28 fold to about 100 fold greater, or is about 28 fold to about 80 fold greater, or is about 28 fold to about 50 fold greater, or is about 28 fold to about 40 fold greater, or is about 28 fold to about 30 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 30 fold to about 100 fold greater, or is about 30 fold to about 80 fold greater, or is about 30 fold to about 50 fold greater, or is about 30 fold to about 40 fold greater, than the effective size of the parental antibody fragment.

In another embodiment, the conjugate has an effective size that is about 40 fold to about 100 fold greater, or is about 40 fold to about 80 fold greater, or is about 40 fold to about 50 fold greater, than the effective size of the parental antibody fragment.

In still another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW of at least about 20,000 D.

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In a further embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW of at least about 30,000 D.

In yet another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW of at least about 40,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW that is at or about 20,000 D to at or about 300,000 D, or is at or about 30,000 D to at or about 300,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW that is at or about 20,000 D to at or about 100,000 D, or is at or about 30,000 D to at or about 100,000 D, or is at or about 40,000 D to at or about 100,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW that is at or about 20,000 D to at or about 70,000 D, or is at or about 30,000 D to at or about 70,000 D, or is at or about 40,000 D to at or about 70,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW that is at or about 20,000 D to at or about 50,000 D, or is at or about 30,000 D to at or about 50,000 D, or is at or about 40,000 D to at or about 50,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one polymer having an actual MW that is at or about 20,000 D to at or about 40,000 D, or is at or about 30,000 D to at or about 40,000 D.

The conjugates of the invention can be made using any suitable technique now known or hereafter developed for derivatizing antibody fragments with polymers. It will be appreciated that the invention is not limited to conjugates utilizing any particular type of linkage between an antibody fragment and a polymer.

The conjugates of the invention include species wherein a polymer is covalently attached to a non-specific site or non-specific sites on the parental antibody fragment, i.e. polymer attachment is not targeted to a particular region or a particular amino acid residue in the parental antibody fragment. In such embodiments, the coupling chemistry can, for example, utilize the free epsilon amino groups of lysine residues in the parental antibody as attachment sites for the polymer, wherein such lysine residue amino groups are randomly derivatized with polymer.

In addition, the conjugates of the invention include species wherein a polymer is covalently attached to a specific site or specific sites on the parental antibody fragment, i.e. polymer attachment is targeted to a particular region or a particular amino acid residue or residues in the parental antibody fragment. In such embodiments, the coupling chemistry can, for example, utilize the free sulfhydryl group of a cysteine residue not in a disulfide bridge in the parental antibody fragment. In one embodiment, one or more cysteine residue(s) is (are) engineered into a selected site or sites in the parental antibody fragment for the purpose of providing a specific attachment site or sites for polymer. The polymer can be activated with any functional group that is capable of reacting specifically with the free sulfhydryl or thiol group(s) on the parental antibody, such as maleimide, sulfhydryl, thiol, triflate, tesylate, aziridine, exirane, and 5-pyridyl

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functional groups. The polymer can be coupled to the parental antibody fragment using any protocol suitable for the chemistry of the coupling system selected, such as the protocols and systems described in Section (II)(1)(b) or in Section (T) of the Examples below.

In another embodiment, polymer attachment is targeted to the hinge region of the parental antibody fragment. The location of the hinge region varies according to the isotype of the parental antibody. Typically, the hinge region of IgG, IgD and IgA isotype heavy chains is contained in a proline rich peptide sequence extending between the C<sub>H</sub>1 and C<sub>H</sub>2 domains. In a preferred embodiment, a cysteine residue or residues is (are) engineered into the hinge region of the parental antibody fragment in order to couple polymer specifically to a selected location in the hinge region.

In one aspect, the invention encompasses a conjugate having any molar ratio of polymer to antibody fragment that endows the conjugate with an apparent size in the desired range as taught herein. The apparent size of the conjugate will depend in part upon the size and shape of the polymer used, the size and shape of the antibody fragment used, the number of polymer molecules attached to the antibody fragment, and the location of such attachment site(s) on the antibody fragment. These parameters can easily be identified and maximized to obtain the a conjugate with the desired apparent size for any type of antibody fragment, polymer and linkage system.

In another aspect, the invention encompasses a conjugate with a polymer to antibody fragment molar ratio of no more than about 10:1, or no more than about 5:1, or no more than about 4:1, or no more than about 3:1, or no more than about 2:1, or no more than 1:1.

In yet another aspect, the invention encompasses a conjugate wherein the antibody fragment is attached to about 10 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In another embodiment, the conjugate contains an antibody fragment attached to about 5 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 4 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 20,000 D, or at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In a further embodiment, the conjugate contains an antibody fragment attached to about 3 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 40,000 D. In an additional embodiment, the conjugate contains an antibody fragment attached to about 2 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 2 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 2 or fewer polymer molecules, each polymer molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. Also provided herein is a conjugate containing an antibody fragment attached to a single polymer molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D.

In still another aspect, the invention encompasses a conjugate wherein every polymer molecule in the c njugate has a molecular weight that is at or about 20,000 D to at or about 300,000 D, or is at or about 300,000 D to at or about 300,000 D, and wherein the

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conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In still another aspect, the invention encompasses a conjugate wherein every polymer molecule in the conjugate has a molecular weight that is at or about 20,000 D to at or about 100,000 D, or is at or about 30,000 D to at or about 100,000 D, or is at or about 40,000 D to at or about 100,000 D, and wherein the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In still another aspect, the invention encompasses a conjugate wherein every polymer molecule in the conjugate has a molecular weight that is at or about 20,000 D to at or about 70,000 D, or is at or about 30,000 D to at or about 70,000 D, or is at or about 40,000 D to at or about 70,000 D, and wherein the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In still another aspect, the invention encompasses a conjugate wherein every polymer molecule in the conjugate has a molecular weight that is at or about 20,000 D to at or about 50,000 D, or is at or about 30,000 D to at or about 50,000 D, or is at or about 40,000 D to at or about 50,000 D, and wherein the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In still another aspect, the invention encompasses a conjugate wherein every polymer molecule in the conjugate has a molecular weight that is at or about 20,000 D to at or about 40,000 D, or is at or about 30,000 D to at or about 40,000 D, and wherein the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

It is believed that the serum half-life, MRT and/or serum clearance rate of any antibody fragment can be greatly improved by derivatizing the antibody fragment with polymer as taught herein. In one embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab. Fab', Fab'-SH, Fv, scFv and F(ab')<sub>2</sub>.

In a preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer

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m lecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In yet another preferred embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In a further embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule and the polymer is coupled to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In an additional embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugate is at least about 20,000 D in molecular weight, or at least about 40,000 D in molecular weight, every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

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In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugat is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

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In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 4 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, every polymer molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, every polymer molecule in the conjugate is attached to the hinge region of the antibody fragment, and the conjugate contains no more than about 10 polymer molecules, or no more than about 5 polymer molecules, or no more than about 3 polymer molecules, or no more than about 2 polymer molecules, or no more than 1 polymer molecule.

In a further embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule in the conjugate is at least about 20,000 D in molecular weight, or at least about 40,000 D in molecular weight, and wherein every polymer molecule in the conjugate is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 30,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, and wherein every polymer molecule in the conjugate is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the

c rresponding cysteine residue in the opposite chain.

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In another embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, and wherein every polymer molecule in the conjugate is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule in the conjugate is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, and wherein every polymer molecule in the conjugate is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, and wherein every polymer molecule in the conjugate is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment attached to no more than about 2 polymer molecules, wherein every polymer molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, and wherein every polymer molecule in the conjugate is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In yet another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at least about 20,000 D in molecular weight, or at least about 30,000 D in molecular weight, or at least about 40,000 D in molecular weight, wherein the polymer

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molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 30,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, wherein the polymer molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, wherein the polymer molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, wherein the polymer molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than I polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, wherein the polymer molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another embodiment, the conjugate contains an antibody fragment selected from the group

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consisting of Fab, Fab', and Fab'-SH, wherein the antib dy fragment is attached to no more than I polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, wherein the polymer molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In still another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at least about 20,000 D in molecular weight, or at least about 30,000 D in molecular weight, or at least about 40,000 D in molecular weight, and wherein the polymer molecule is attached to the hinge region of the antibody fragment.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 30,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, and wherein the polymer molecule is attached to the hinge region of the antibody fragment.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, and wherein the polymer molecule is attached to the hinge region of the antibody fragment.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, and wherein the polymer molecule is attached to the hinge region of the antibody fragment.

In another embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, and wherein the polymer molecule is attached to the hinge region of the antibody fragment.

In another embodiment, the conjugate contains an antibody fragment selected from the group

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consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 polymer molecule, wherein the polymer molecule is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, and wherein the polymer molecule is attached to the hinge region of the antibody fragment.

Although any type of polymer is contemplated for use in constructing the conjugates of the invention, including the polymers and chemical linkage systems described in Section (II)(1)(b) below, polyethylene glycol (PEG) polymers are preferred for use herein.

In one embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW of at least about 20,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW of at least about 30,000 D.

In yet another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW of at least about 40,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW that is at or about 20,000 D to at or about 300,000 D, or is at or about 30,000 D to at or about 300,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW that is at or about 20,000 D to at or about 100,000 D, or is at or about 30,000 D to at or about 100,000 D, or is at or about 40,000 D to at or about 100,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW that is at or about 20,000 D to at or about 70,000 D, or is at or about 30,000 D to at or about 70,000 D, or is at or about 40,000 D to at or about 70,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW that is at or about 20,000 D to at or about 50,000 D, or is at or about 30,000 D to at or about 50,000 D, or is at or about 40,000 D to at or about 50,000 D.

In another embodiment, the conjugate is an antibody fragment covalently attached to at least one PEG having an actual MW that is at or about 20,000 D to at or about 40,000 D, or is at or about 30,000 D to at or about 40,000 D.

In another aspect, the invention encompasses a conjugate with a PEG to antibody fragment molar ratio of no more than about 10:1, or no more than about 5:1, or no more than about 4:1, or no more than about 3:1, or no more than about 2:1, or no more than 1:1.

In yet another aspect, the invention encompasses a conjugate wherein the antibody fragment is attached to about 10 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In another embodiment, the conjugate contains an antibody fragment attached to about 5 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In still another embodiment, the conjugate contains an antibody fragment attached to about 4 or fewer PEG

molecules, each PEG molecule having a molecular weight of at least ab ut 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In a further embodiment, the conjugate contains an antibody fragment attached to about 3 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In an additional embodiment, the conjugate contains an antibody fragment attached to about 2 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. Also provided herein is a conjugate containing an antibody fragment attached to a single PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D.

In another aspect, the invention encompasses a conjugate wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 300,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules. or no more than about 2 PEG molecules, or no more than 1 PEG molecules.

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In another aspect, the invention encompasses a conjugate wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecules.

In another aspect, the invention encompasses a conjugate wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another aspect, the invention encompasses a conjugate wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecules.

In another aspect, the invention encompasses a conjugate wherein the antibody fragment is

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derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In still another aspect, the invention encompasses a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', Fab'-SH and F(ab')2, wherein the antibody fragment is attached to about 10 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In another embodiment, the foregoing conjugate contains an antibody fragment attached to about 5 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In still another embodiment, the foregoing conjugate contains an antibody fragment attached to about 4 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In a further embodiment, the foregoing conjugate contains an antibody fragment attached to about 3 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. In an additional embodiment, the foregoing conjugate contains an antibody fragment attached to about 2 or fewer PEG molecules, each PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. Also provided herein is the foregoing conjugate that contains an antibody fragment attached to a single PEG molecule having a molecular weight of at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D.

In another aspect, the invention encompasses a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', Fab'-SH and F(ab')<sub>2</sub>, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 300,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another aspect, the invention encompasses a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', Fab'-SH and F(ab')<sub>2</sub>, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG

molecule.

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In another aspect, the invention encompasses a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', Fab'-SH and F(ab')<sub>2</sub>, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another aspect, the invention encompasses a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', Fab'-SH and F(ab')<sub>2</sub>, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another aspect, the invention encompasses a conjugate containing an antibody fragment selected from the group consisting of Fab. Fab', Fab'-SH and F(ab')<sub>2</sub>, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In a preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG having a molecular weight of at least about 20,000D, or at least about 30,000D, or at least about 40,000D, and wherein every PEG molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG having a molecular weight that is at or about 20,000 D to about 300,000 D, or is at or about 30,000 D to at r about 300,000 D, or is at or about 40,000 D to at or about 300,000 D, and wherein every PEG molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG having a molecular weight that is at or about 20,000 D to about 100,000 D, or is at or about 30,000 D to at or about

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100,000 D, or is at or about 40,000 D to at or about 100,000 D, and wherein every PEG molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG having a molecular weight that is at or about 20,000 D to about 70,000 D, or is at or about 30,000 D to at or about 70,000 D, or is at or about 40,000 D to at or about 70,000 D, and wherein every PEG molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG having a molecular weight that is at or about 20,000 D to about 50,000 D, or is at or about 30,000 D to at or about 50,000 D, and wherein every PEG molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG having a molecular weight that is at or about 20,000 D to about 40,000 D, or is at or about 30,000 D to at or about 40,000 D, and wherein every PEG molecule in the conjugate is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at least about 20,000D in molecular weight, or at least about 30,000D in molecular weight, or at least about 40,000D in molecular weight, wherein every PEG molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 300,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, wherein every PEG molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular

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weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, wherein every PEG molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, wherein every PEG molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, wherein every PEG molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 4 PEG molecules, or no more than about 3 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, wherein every PEG molecule in the conjugate molecule is attached to the hinge region of the antibody fragment, and wherein the conjugate contains no more than about 10 PEG molecules, or no more than about 5 PEG molecules, or no more than about 2 PEG molecules, or no more than about 2 PEG molecules, or no more than 1 PEG molecule.

In yet another preferred embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment derivatized with PEG, wherein every PEG molecule in the conjugate is at least about 20,000D in molecular weight, or at least about 30,000D in molecular weight, or at least about 40,000D in molecular weight, wherein the antibody fragment is attached to no more than about 2 PEG molecules, and wherein every PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would

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ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the c rresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 30,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, wherein the antibody fragment is attached to no more than about 2 PEG molecules, and wherein every PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, or is at or about 40,000 D to at or about 100,000 D in molecular weight, wherein the antibody fragment is attached to no more than about 2 PEG molecules, and wherein every PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, wherein the antibody fragment is attached to no more than about 2 PEG molecules, and wherein every PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains a F(ab')<sub>2</sub> antibody fragment derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, wherein the antibody fragment is attached to no more than about 2 PEG molecules, and wherein every PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains a F(ab')2 antibody fragment derivatized

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with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in m lecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, wherein the antibody fragment is attached to no more than about 2 PEG molecules, and wherein every PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In still another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at least about 20,000 D in molecular weight, or at least about 30,000 in molecular weight, or at least about 40,000 D in molecular weight, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 300,000 D in molecular weight, or is at or about 30,000 D to at or about 300,000 D in molecular weight, or is at or about 40,000 D to at or about 300,000 D in molecular weight, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 100,000 D in molecular weight, or is at or about 30,000 D to at or about 100,000 D in molecular weight, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 70,000 D in molecular weight, or is at or about 30,000 D to at or about 70,000 D in molecular weight, or is at or about 40,000 D to at or about 70,000 D in molecular weight, wherein the antibody fragment is attached to no more than 1 PEG

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molecule, and wherein the PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 50,000 D in molecular weight, or is at or about 30,000 D to at or about 50,000 D in molecular weight, or is at or about 40,000 D to at or about 50,000 D in molecular weight, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is derivatized with PEG, wherein every PEG molecule in the conjugate is at or about 20,000 D to at or about 40,000 D in molecular weight, or is at or about 30,000 D to at or about 40,000 D in molecular weight, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is attached to a cysteine residue in the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains, wherein the disulfide bridge is avoided by substituting another amino acid, such as serine, for the corresponding cysteine residue in the opposite chain.

It will be appreciated that all of the above-described embodiments of the invention utilizing PEG polymers include conjugates wherein the PEG polymer(s) is (are) linear or branched. In a preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is branched and at least about 40,000 D in molecular weight. In a particularly surprising and unexpected finding, the inventors discovered that the foregoing conjugate exhibits a serum half-life, MRT and serum clearance rate approaching that of full length antibody as shown in Example X below.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than I PEG molecule, and wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 300,000 D.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than I PEG molecule, and wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 100,000 D.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the

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group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no m re than 1 PEG molecule, and wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 70,000 D.

In another preferred embodiment, the conjugate contains an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, and wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 50,000 D.

In another preferred embodiment, the invention provides a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, wherein the PEG molecule is branched and at least 40,000D in molecular weight, and the PEG molecule is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the invention provides a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 300,000 D, and the PEG molecule is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the invention provides a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 100,000 D, and the PEG molecule is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the invention provides a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 70,000 D, and the PEG molecule is attached to the hinge region of the antibody fragment.

In another preferred embodiment, the invention provides a conjugate containing an antibody fragment selected from the group consisting of Fab, Fab', and Fab'-SH, wherein the antibody fragment is attached to no more than 1 PEG molecule, wherein the PEG molecule is branched and has a molecular weight that is at or about 40,000 D to at or about 50,000 D, and the PEG molecule is attached to the hinge region of the antibody fragment.

In one aspect, the invention provides any of the above-described conjugates wherein the c njugate contains no more than one antibody fragment. Additionally provided herein is any of the above-described conjugates wherein the conjugate contains one or more antibody fragment(s) covalently linked to one or more polymer molecule(s), such as conjugates containing two or more antibody fragments covalently linked together by polymer molecule(s). In one embodiment, a polymer molecule is used to link together two antibody fragments to form a dumbbell-shaped structure. Also encompassed herein are conjugates formed

by more than two antibody fragments joined by polymer molecule(s) to form a rosette or other shapes. The antibody fragments in such structures can be of the same or different fragment type and can have the same antigen specificity or have different antigen specificities. Such structures can be made by using a polymer molecule derivatized with multiple functional groups permitting the direct attachment, or the attachment by means of bi- or multi-functional linkers, of two or more antibody fragments to the polymer backbone.

In another aspect, the invention encompasses any of the above-described conjugates utilizing an antibody fragment comprising an antigen recognition site that binds to rabbit IL-8 and/or human IL-8. In yet another aspect, the invention encompasses any of the above-described conjugates utilizing an antibody fragment comprising 6G4.2.5LV/L1N35A or 6G4.2.5LV/L1N35E as defined below. In still another aspect, the invention encompasses any of the above-described conjugates utilizing an antibody fragment comprising 6G4.5.2.5HV11 as defined below. In a further aspect, the invention encompasses any of the abovedescribed conjugates utilizing an antibody fragment comprising hu6G4.2.5LV/L1N35A or hu6G4.2.5LV/L1N35E as defined below. In an additional aspect, the invention encompasses any of the above-described conjugates utilizing an antibody fragment comprising hu6G4.2.5HV. Further encompassed herein are any of the above-described conjugates utilizing an antibody fragment comprising 6G4.2.5LV/L1N35A or 6G4.2.5LV/L1N35E and further comprising the CDRs of 6G4.2.5HV as defined below. Also encompassed herein are any of the above described conjugates utilizing an antibody fragment comprising hu6G4.2.5LV/L1N35A or hu6G4.2.5LV/L1N35E and further comprising hu6G4.2.5HV as defined below. Additionally encompassed herein are any of the above-described conjugates utilizing an antibody fragment comprising 6G4.2.5LV11N35A or 6G4.2.5LV11N35E as defined below. Further provided herein are any of the above-described conjugates utilizing an antibody fragment comprising 6G4.2.5LV11N35A or 6G4.2.5LV11N35E and further comprising 6G4.2.5HV11 as defined below.

## a. Production of Antibody Fragments

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Antibody fragments can be produced by any method known in the art. Generally, an antibody fragment is derived from a parental intact antibody. The parental antibody can be generated by raising polyclonal sera against the desired antigen by multiple subcutaneous (sc) or intraperitoneal (ip) injections of antigen and an adjuvant, such as monophosphoryl lipid A (MPL)/trehalose dicrynomycolate (TDM) (Ribi Immunochem. Research, Inc., Hamilton, MT), at multiple sites. Two weeks later the animals are boosted. 7 to 14 days later animals are bled and the serum is assayed for anti-antigen titer. Animals are boosted until titer plateaus. Sera are harvested from animals, and polyclonal antibodies are isolated from sera by conventional immunoglobulin purification procedures, such as protein A-Sepharose chromatography, hydroxylapatite chromatography, gel filtration, dialysis, or antigen affinity chromatography. The desired antibody fragments can be generated from purified polyclonal antibody preparations by conventional enzymatic methods, e.g. F(ab')<sub>2</sub> fragments are produced by pepsin cleavage of intact antibody, and Fab fragments are produced by briefly digesting intact antibody with papain.

Alternatively, antibody fragments are derived from monoclonal antibodies generated against the desired antigen. Monoclonal antibodies may be made using the hybridoma method first described by Kohler

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et al., Nature, 256:495 (1975), or may be made by recombinant DNA methods (U.S. Patent No. 4,816,567).

In the hybridoma method, a mouse or other appropriate host animal, such as a hamster or macaque monkey, is immunized as hereinabove described to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the protein used for immunization. Alternatively, lymphocytes may be immunized *in vitro*. Lymphocytes then are fused with myeloma cells using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, *Monoclonal Antibodies: Principles and Practice*, pp.59-103 (Academic Press, 1986)).

The hybridoma cells thus prepared are seeded and grown in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, parental myeloma cells. For example, if the parental myeloma cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine (HAT medium), which substances prevent the growth of HGPRT-deficient cells.

Preferred myeloma cells are those that fuse efficiently, support stable high-level production of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. Among these, preferred myeloma cell lines are murine myeloma lines, such as those derived from MOP-21 and M.C.-11 mouse tumors available from the Salk Institute Cell Distribution Center, San Diego, California USA, and SP-2 or X63-Ag8-653 cells available from the American Type Culture Collection, Rockville, Maryland USA. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); Brodeur et al., Monoclonal Antibody Production Techniques and Applications, pp. 51-63 (Marcel Dekker, Inc., New York, 1987)).

Culture medium in which hybridoma cells are growing is assayed for production of monoclonal antibodies directed against the antigen. Preferably, the binding specificity of monoclonal antibodies produced by hybridoma cells is determined by immunoprecipitation or by an *in vitro* binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA).

The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson et al., Anal. Biochem., 107:220 (1980).

After hybridoma cells are identified that produce antibodies of the desired specificity, affinity, and/or activity, the clones may be subcloned by limiting dilution procedures and grown by standard methods (Goding, Monoclonal Antibodies: Principles and Practice, pp.59-103 (Academic Press, 1986)). Suitable culture media for this purpose include, for example, D-MEM or RPMI-1640 medium. In addition, the hybridoma cells may be grown in vivo as ascites tumors in an animal.

The monoclonal antibodies secreted by the subclones are suitably separated from the culture medium, ascites fluid, or serum by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

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DNA encoding the monoclonal antibodies is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide pr bes that are capable of binding specifically to genes encoding the heavy and light chains of the monoclonal antibodies). The hybridoma cells serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as E. coli cells, simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. Review articles on recombinant expression in bacteria of antibody-encoding DNA include Skerra et al., Curr. Opinion in Immunol., 5: 256 (1993) and Pluckthun, Immunol. Revs., 130: 151 (1992).

In a preferred embodiment, the antibody fragment is derived from a humanized antibody. Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. It will be appreciated that variable domain sequences obtained from any non-human animal phage display library-derived Fv clone or from any non-human animal hybridoma-derived antibody clone provided as described herein can serve as the "import" variable domain used in the construction of the humanized antibodies of the invention. Humanization can be essentially performed following the method of Winter and co-workers (Jones et al., Nature, 321: 522 (1986): Riechmann et al., Nature, 332: 323 (1988); Verhoeyen et al., Science, 239: 1534 (1988)), by substituting non-human animal, e.g. rodent, CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (Cabilly et al., supra), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in non-human animal, e.g. rodent, antibodies.

The choice of human variable domains, both light and heavy, to be used in making the humanized antibodies is very important to reduce antigenicity. According to the so-called "best-fit" method, the sequence of the variable domain of a non-human animal, e.g. rodent, antibody is screened against the entire library of known human variable-domain sequences. The human sequence which is closest to that of the non-human animal is then accepted as the human framework (FR) for the humanized antibody (Sims et al., J. Immunol., 151: 2296 (1993); Chothia and Lesk, J. Mol. Biol., 196: 901 (1987)). Another method uses a particular framework derived from the consensus sequence of all human antibodies of a particular subgroup light or heavy chains. The same framework can be used for several different humanized antibodies (Carter et al., Proc. Natl. Acad. Sci USA, 89: 4285 (1992); Presta et al., J. Immunol., 151: 2623 (1993)).

It is also important that antibodies be humanized with retention of high affinity for the antigen and other favorable biological properties. To achieve this goal, according to a preferred method, humanized antibodies are prepared by a process of analysis of the parental sequences and various c nceptual humanized products using three-dimensional models of the parental and humanized sequences. Three-dimensional

immunoglobulin models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influence the ability of the candidate immunoglobulin to bind to its antigen. In this way, FR residues can be selected and combined from the consensus and import sequences so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved. In general, the CDR residues are directly and most substantially involved in influencing antigen binding.

In addition, antibody fragments for use herein can be derived from human monoclonal antibodies. Human monoclonal antibodies against the antigen of interest can be made by the hybridoma method. Human myeloma and mouse-human heteromyeloma cell lines for the production of human monoclonal antibodies have been described, for example, by Kozbor J. Immunol., 133: 3001 (1984); Brodeur et al., Monoclonal Antibody Production Techniques and Applications, pp. 51-63 (Marcel Dekker, Inc., New York, 1987); and Boerner et al., J. Immunol., 147: 86 (1991).

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It is now possible to produce transgenic animals (e.g. mice) that are capable, upon immunization, of producing a full repertoire of human antibodies in the absence of endogenous immunoglobulin production. For example, it has been described that the homozygous deletion of the antibody heavy-chain joining region (JH) gene in chimeric and germ-line mutant mice results in complete inhibition of endogenous antibody production. Transfer of the human germ-line immunoglobulin gene array in such germ-line mutant mice will result in the production of human antibodies upon antigen challenge. See, e.g., Jakobovits et al., Proc. Natl. Acad. Sci USA, 90: 2551 (1993); Jakobovits et al., Nature, 362: 255 (1993); Bruggermann et al., Year in Immunol., 7: 33 (1993).

Alternatively, phage display technology (McCafferty et al., Nature 348:552 (1990)) can be used to produce human antibodies and antibody fragments in vitro, from immunoglobulin variable (V) domain gene repertoires from unimmunized donors. According to this technique, antibody V domain genes are cloned inframe into either a major or minor coat protein gene of a filamentous bacteriophage, such as M13 or fd, and displayed as functional antibody fragments on the surface of the phage particle. Because the filamentous particle contains a single-stranded DNA copy of the phage genome, selections based on the functional properties of the antibody also result in selection of the gene encoding the antibody exhibiting those properties. Thus, the phage mimics some of the properties of the B-cell. Phage display can be performed in a variety of formats; for their review see, e.g., Johnson et al., Current Opinion in Structural Biology 3:564 (1993). Several sources of V-gene segments can be used for phage display. Clackson et al., Nature 352:624 (1991) isolated a diverse array of anti-oxazolone antibodies from a small random combinatorial library of V genes derived from the spleens of immunized mice. A repertoire of V genes from unimmunized human donors can be constructed and antibodies to a diverse array of antigens (including self-antigens) can be isolated essentially following the techniques described by Marks et al., J. Mol. Biol. 222:581 (1991), or Griffith et al., EMBO J. 12:725 (1993).

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In a natural immune response, antibody genes accumulate mutations at a high rate (somatic hypermutation). Some of the changes introduced will confer higher affinity, and B cells displaying high-affinity surface immunoglobulin are preferentially replicated and differentiated during subsequent antigen challenge. This natural process can be mimicked by employing the technique known as "chain shuffling" (Marks et al., Bio/Technol. 10:779 (1992)). In this method, the affinity of "primary" human antibodies obtained by phage display can be improved by sequentially replacing the heavy and light chain V region genes with repertoires of naturally occurring variants (repertoires) of V domain genes obtained from unimmunized donors. This technique allows the production of antibodies and antibody fragments with affinities in the nM range. A strategy for making very large phage antibody repertoires has been described by Waterhouse et al., Nucl. Acids Res. 21:2265 (1993).

Gene shuffling can also be used to derive human antibodies from non-human, e.g. rodent, antibodies, where the human antibody has similar affinities and specificities to the starting non-human antibody. According to this method, which is also called "epitope imprinting", either the heavy or light chain variable region of a non-human antibody fragment obtained by phage display techniques as described above is replaced with a repertoire of human V domain genes, creating a population of non-human chain/human chain scFv or Fab chimeras. Selection with antigen results in isolation of a non-human chain/human chain chimeric scFv or Fab wherein the human chain restores the antigen binding site destroyed upon removal of the corresponding non-human chain in the primary phage display clone, i.e. the epitope governs (imprints) the choice of the human chain partner. When the process is repeated in order to replace the remaining non-human chain, a human antibody is obtained (see PCT WO 93/06213 published April 1, 1993). Unlike traditional humanization of non-human antibodies by CDR grafting, this technique provides completely human antibodies, which have no FR or CDR residues of non-human origin.

The invention also encompasses the use of bispecific and heteroconjugate antibody fragments having specificities for at least two different antigens. Bispecific and heteroconjugate antibodies can be prepared as full length antibodies or as antibody fragments (e.g. F(ab')<sub>2</sub> bispecific antibody fragments). Antibody fragments having more than two valencies (e.g. trivalent or higher valency antibody fragments) are also contemplated for use herein. Bispecific antibodies, heteroconjugate antibodies, and multi-valent antibodies can be prepared as described in Section (II)(3)(C) below.

As described above, DNA encoding the monoclonal antibody or antibody fragment of interest can be isolated from its hybridoma or phage display clone of origin, and then manipulated to create humanized and/or affinity matured constructs. In addition, known techniques can be employed to introduce an amino acid residue or residues into any desired location on the polypeptide backbone of the antibody fragment, e.g. a cysteine residue placed in the hinge region of the heavy chain, thereby providing a site for specific attachment of polymer molecule(s). In one embodiment, the native cysteine residue in either the light or heavy chain of the antibody fragment that would ordinarily form the disulfide bridge linking the light and heavy chains is substituted with another amino acid, such as serine, in order to leave the partner cysteine residue in the opposite chain with a free suflhydryl for specific attachment of polymer molecule.

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Upon construction of the desired antibody or antibody fragment-encoding clone, the clone can be used for recombinant production of the antibody fragment as described in Section (II)(4) below. Finally, the antibody or antibody fragment product can be recovered from host cell culture and purified as described in Section (II)(4)(F) below. In the case of embodiments utilizing an antibody fragment engineered to lack a cysteine residue that ordinarily forms the disulfide bridge between the light and heavy chains as described above, preferred recombinant production systems include bacterial expression and product recovery procedures utilizing the low pH osmotic shock method described in the "Alternative Fab'-SH Purification" section of Example T below. If a full length antibody is produced, the desired antibody fragment can be obtained therefrom by subjecting the intact antibody to enzymatic digestion according to known methods, e.g. as described in Section (II)(4)(G) below.

# b. Construction of Antibody Fragment-Polymer Conjugates

The antibody fragment-polymer conjugates of the invention can be made by derivatizing the desired antibody fragment with an inert polymer. It will be appreciated that any inert polymer which provides the conjugate with the desired apparent size or which has the selected actual MW as taught herein is suitable for use in constructing the antibody fragment-polymer conjugates of the invention.

Many inert polymers are suitable for use in pharmaceuticals. See, e.g., Davis et al., Biomedical Polymers: Polymeric Materials and Pharmaceuticals for Biomedical Use, pp.441-451 (1980). embodiments of the invention, a non-proteinaceous polymer is used. The nonproteinaceous polymer ordinarily is a hydrophilic synthetic polymer, i.e., a polymer not otherwise found in nature. However, polymers which exist in nature and are produced by recombinant or in vitro methods are also useful, as are polymers which are isolated from native sources. Hydrophilic polyvinyl polymers fall within the scope of this invention, e.g. polyvinylalcohol and polyvinylpyrrolidone. Particularly useful are polyalkylene ethers such as polyethylene glycol (PEG); polyoxyalkylenes such as polyoxyethylene, polyoxypropylene, and block copolymers of polyoxyethylene and polyoxypropylene (Pluronics); polymethacrylates; carbomers; branched or unbranched polysaccharides which comprise the saccharide monomers D-mannose, D- and Lgalactose, fucose, fructose, D-xylose, L-arabinose, D-glucuronic acid, sialic acid, D-galacturonic acid, Dmannuronic acid (e.g. polymannuronic acid, or alginic acid), D-glucosamine, D-galactosamine, D-glucose and neuraminic acid including homopolysaccharides and heteropolysaccharides such as lactose, amylopectin, starch, hydroxyethyl starch, amylose, dextrane sulfate, dextran, dextrins, glycogen, or the polysaccharide subunit of acid mucopolysaccharides, e.g. hyaluronic acid; polymers of sugar alcohols such as polysorbitol and polymannitol; heparin or heparon. The polymer prior to cross-linking need not be, but preferably is, water soluble, but the final conjugate must be water soluble. Preferably, the conjugate exhibits a water solubility of at least about 0.01 mg/ml, and more preferably at least about 0.1 mg/ml, and still more preferably at least about 1 mg/ml. In addition, the polymer should not be highly immunogenic in the conjugate form, nor should it possess viscosity that is incompatible with intravenous infusion or injection if the conjugate is intended to be administered by such routes.

In one embodiment, the polymer contains only a single group which is reactive. This helps to

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avoid cross-linking of protein molecules. However, it is within the scope herein to maximize reaction conditions to reduce cross-linking, or to purify the reaction products through gel filtration or ion exchange chromatography to recover substantially homogenous derivatives. In other embodiments, the polymer contains two or more reactive groups for the purpose of linking multiple antibody fragments to the polymer backbone. Again, gel filtration or ion exchange chromatography can be used to recover the desired derivative in substantially homogeneous form.

The molecular weight of the polymer can range up to about 500,000 D, and preferably is at least about 20,000 D, or at least about 30,000 D, or at least about 40,000 D. The molecular weight chosen can depend upon the effective size of the conjugate to be achieved, the nature (e.g. structure, such as linear or branched) of the polymer, and the degree of derivatization, i.e. the number of polymer molecules per antibody fragment, and the polymer attachment site or sites on the antibody fragment.

The polymer can be covalently linked to the antibody fragment through a multifunctional crosslinking agent which reacts with the polymer and one or more amino acid residues of the antibody fragment to be linked. However, it is also within the scope of the invention to directly crosslink the polymer by reacting a derivatized polymer with the antibody fragment, or vice versa.

The covalent crosslinking site on the antibody fragment includes the N-terminal amino group and epsilon amino groups found on lysine residues, as well as other amino, imino, carboxyl, sulfhydryl, hydroxyl or other hydrophilic groups. The polymer may be covalently bonded directly to the antibody fragment without the use of a multifunctional (ordinarily bifunctional) crosslinking agent. Covalent binding to amino groups is accomplished by known chemistries based upon cyanuric chloride, carbonyl diimidazole, aldehyde reactive groups (PEG alkoxide plus diethyl acetal of bromoacetaldehyde; PEG plus DMSO and acetic anhydride, or PEG chloride plus the phenoxide of 4-hydroxybenzaldehyde, activated succinimidyl esters, activated dithiocarbonate PEG, 2,4,5-trichlorophenylcloroformate or P-nitrophenylcloroformate activated PEG.) Carboxyl groups are derivatized by coupling PEG-amine using carbodiimide. Sulfhydryl groups are derivatized by coupling to maleimido-substituted PEG (e.g. alkoxy-PEG amine plus sulfosuccinimidyl 4-(N-maleimidomethyl)cyclohexane-1-carboxylate) as described in WO 97/10847 published March 27, 1997, or PEG-maleimide commercially available from Shearwater Polymers, Inc., Huntsville, AL). Alternatively, free amino groups on the antibody fragment (e.g. epsilon amino groups on lysine residues) can be thiolated with 2-imino-thiolane (Traut's reagent) and then coupled to maleimide-containing derivatives of PEG as described in Pedley et al., Br. J. Cancer, 70: 1126-1130 (1994).

The polymer will bear a group which is directly reactive with an amino acid side chain, or the N- or C-terminus of the polypeptide linked, or which is reactive with the multifunctional cross-linking agent. In general, polymers bearing such reactive groups are known for the preparation of immobilized proteins. In order to use such chemistries here, one should employ a water soluble polymer otherwise derivatized in the same fashion as insoluble polymers heretofore employed for protein immobilization. Cyanogen bromide activation is a particularly useful procedure to employ in crosslinking polysaccharides.

"Water soluble" in reference to the starting polymer means that the polymer or its reactive

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intermediate used for conjugation is sufficiently water soluble to participate in a derivatization reaction.

The degree of substitution with such a polymer will vary depending upon the number of reactive sites on the antibody fragment, the molecular weight, hydrophilicity and other characteristics of the polymer, and the particular antibody fragment derivatization sites chosen. In general, the conjugate contains from 1 to about 10 polymer molecules, but greater numbers of polymer molecules attached to the antibody fragments of the invention are also contemplated. The desired amount of derivatization is easily achieved by using an experimental matrix in which the time, temperature and other reaction conditions are varied to change the degree of substitution, after which the level of polymer substitution of the conjugates is determined by size exclusion chromatography or other means known in the art.

The polymer, e.g. PEG, is cross-linked to the antibody fragment by a wide variety of methods known per se for the covalent modification of proteins with nonproteinaceous polymers such as PEG. Certain of these methods, however, are not preferred for the purposes herein. Cyanuronic chloride chemistry leads to many side reactions, including protein cross-linking. In addition, it may be particularly likely to lead to inactivation of proteins containing sulfhydryl groups. Carbonyl diimidazole chemistry (Beauchamp et al., Anal Biochem. 131, 25-33 [1983]) requires high pH (>8.5), which can inactivate proteins. Moreover, since the "activated PEG" intermediate can react with water, a very large molar excess of "activated PEG" over protein is required. The high concentrations of PEG required for the carbonyl diimidazole chemistry also led to problems in purification, as both gel filtration chromatography and hydrophilic interaction chromatography are adversely affected. In addition, the high concentrations of "activated PEG" may precipitate protein, a problem that per se has been noted previously (Davis, U.S. Patent No. 4,179,337). On the other hand, aldehyde chemistry (Royer, U.S. Patent No. 4,002,531) is more efficient since it requires only a 40-fold molar excess of PEG and a 1-2 hr incubation. However, the manganese dioxide suggested by Royer for preparation of the PEG aldehyde is problematic "because of the pronounced tendency of PEG to form complexes with metal-based oxidizing agents" (Harris et al., J. Polym. Sci. Polym. Chem. Ed. 22, 341-52 [1984]). The use of a Moffatt oxidation, utilizing DMSO and acetic anhydride, obviates this problem. In addition, the sodium borohydride suggested by Royer must be used at high pH and has a significant tendency to reduce disulfide bonds. In contrast, sodium cyanoborohydride, which is effective at neutral pH and has very little tendency to reduce disulfide bonds is preferred. In another preferred embodiment, maleimido-activated PEG is used for coupling to free thiols on the antibody fragment.

Functionalized PEG polymers to modify the antibody fragments of the invention are available from Shearwater Polymers, Inc. (Huntsville, AL). Such commercially available PEG derivatives include, but are not limited to, amino-PEG, PEG amino acid esters, PEG-hydrazide, PEG-thiol, PEG-succinate, carboxymethylated PEG, PEG-propionic acid, PEG amino acids, PEG succinimidyl succinate, PEG succinimidyl propionate, succinimidyl ester of carboxymethylated PEG, succinimidyl carbonate of PEG, succinimidyl esters of amino acid PEGs, PEG-oxycarbonylimidazole, PEG-nitrophenyl carbonate, PEG tresylate, PEG-glycidyl ether, PEG-aldehyde, PEG vinylsulfone, PEG-maleimide, PEG-orthopyridyl-

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disulfide, heterofunctional PEGs, PEG vinyl derivatives, PEG silanes, and PEG phospholides. The reaction conditions for coupling these PEG derivatives will vary depending on the protein, the desired degree of PEGylation, and the PEG derivative utilized. Some factors involved in the choice of PEG derivatives include: the desired point of attachment (such as lysine or cysteine R-groups), hydrolytic stability and reactivity of the derivatives, stability, toxicity and antigenicity of the linkage, suitability for analysis, etc. Specific instructions for the use of any particular derivative are available from the manufacturer.

The conjugates of this invention are separated from the unreacted starting materials by gel filtration or ion exchange HPLC. Heterologous species of the conjugates are purified from one another in the same fashion.

The conjugates may also be purified by ion-exchange chromatography. The chemistry of many of the electrophilically activated PEG's results in a reduction of amino group charge of the PEGylated product. Thus, high resolution ion exchange chromatography can be used to separate the free and conjugated proteins, and to resolve species with different levels of PEGylation. In fact, the resolution of different species (e.g. containing one or two PEG residues) is also possible due to the difference in the ionic properties of the unreacted amino acids. In one embodiment, species with difference levels of PEGylation are resolved according to the methods described in WO 96/34015 (International Application No. PCT/US96/05550 published October 31, 1996).

In a preferred embodiment, the conjugate is generated by utilizing the derivatization and purification methods described in Section (T) of the Examples below.

In one aspect, the invention provides any of the above-described conjugates formed by its component parts, i.e. one or more antibody fragment(s) covalently attached to one or more polymer molecule(s), without any extraneous matter in the covalent molecular structure of the conjugate.

#### c. Other Derivatives of Large Effective Size Conjugates

In another aspect, any of the above-described conjugates can be modified to contain one or more component(s) in addition to the antibody fragment component(s) and polymer component(s) that form the conjugate, wherein the modification does not alter the essential functional property of the conjugate, namely, the substantially improved serum half-life, MRT and/or serum clearance rate as compared to that of the parental antibody fragment from which the conjugate is derived. In one embodiment, the invention provides any of the above-described conjugates modified to incorporate one or more nonproteinaceous functional group(s). For example, the conjugate can be modified to incorporate nonproteinaceous labels or reporter molecules, such as radiolabels, including any radioactive substance used in medical treatment or imaging or used as an effector function or tracer in an animal model, such as radioisotopic labels <sup>99</sup>Tc, <sup>90</sup>Y, <sup>111</sup>In, <sup>32</sup>P, <sup>14</sup>C, <sup>125</sup>I, <sup>3</sup>H, <sup>131</sup>I, <sup>11</sup>C, <sup>15</sup>O, <sup>13</sup>N, <sup>18</sup>F, <sup>35</sup>S, <sup>51</sup>Cr, <sup>57</sup>To, <sup>226</sup>Ra, <sup>60</sup>Co, <sup>59</sup>Fe, <sup>75</sup>Se, <sup>152</sup>Eu, <sup>67</sup>Cu, <sup>217</sup>Ci, <sup>211</sup>At, <sup>212</sup>Pb, <sup>47</sup>Sc, <sup>109</sup>Pd, <sup>234</sup>Th, <sup>40</sup>K, and the like, non-radioisotopic labels such as <sup>157</sup>Gd, <sup>55</sup>Mn, <sup>52</sup>Tr, <sup>56</sup>Fe, etc., fluroescent or chemiluminescent labels, including fluorophores such as rare earth chelates, fluorescein and its derivatives, rhodamine and its derivatives, isothiocyanate, phycoerythrin, phycocyanin,

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all phycocyanin, o-phthaladehyde, fluorescamine, <sup>152</sup>Eu, dansyl, umbelliferone, luciferin, luminal label, isoluminal label, an aromatic acridinium ester label, an imidazole label, an acridimium salt label, an oxalate ester label, an aequorin label, 2,3-dihydrophthalazinediones, biotin/avidin, spin labels, stable free radicals, and the like.

Conventional methods are available to bind these labels covalently to the polypeptide antibody fragment or polymer component of the conjugate. In one aspect, any conjugate of the invention is modified by derivatizing the antibody fragment component with any of the above-described non-proteinaceous labels, wherein the label is directly or indirectly (through a coupling agent) attached to the antibody fragment, and wherein such derivatization of the antibody fragment does not contribute or introduce any polymer moiety into the molecular structure of the conjugate. For instance, coupling agents such as dialdehydes, carbodiimides, dimaleimides, bis-imidates, bis-diazotized benzidine, and the like can be used to tag the antibody fragment with the above-described fluorescent or chemiluminescent labels. See, for example, U.S. Pat. No. 3,940,475 (fluorimetry), Morrison, Meth. Enzymol., 32b, 103 (1974), Svyanen et al., J. Biol. Chem., 284, 3762 (1973), and Bolton and Hunter, Biochem. J., 133, 529 (1973).

In the case of embodiments utilizing radiolabels, both direct and indirect labeling can be used to incorporate the selected radionuclide into the conjugate. As used herein in the context of radiolabeling, the phrases "indirect labeling" and "indirect labeling approach" both mean that a chelating agent is covalently attached to the antibody fragment moiety or polymer moiety of the conjugate and at least one raidonuclide is inserted into the chelating agent. Preferred chelating agents and radionuclides are set forth in Srivagtava, S.C. and Mease, R.C., "Progress in Research on Ligands, Nuclides and Techniques for Labeling Monoclonal Antibodies," Nucl. Med. Bio., 18(6): 589-603 (1991). A particularly preferred chelating agent is 1isothiocycmatobenzyl-3-methyldiothelene triaminepent acetic acid ("MX-DTPA"). As used herein in the context of radiolabeling, the phrases "direct labeling" and "direct labeling approach" both mean that a radionuclide is covalently attached directly to the antibody fragment moiety (typically via an amino acid residue) or to the polymer moiety of the conjugate. Preferred radionuclides for use in direct labeling of conjugate are provided in Srivagtava and Mease, supra. In one embodiment, the conjugate is directly labeled with 131 covalently attached to tyrosine residues. In another embodiment, the antibody fragment component of the conjugate is directly or indirectly labeled with any of the above-described radiolabels, wherein such labeling of the antibody fragment does not contribute or introduce any polymer moiety into the molecular structure of the conjugate.

# d. Therapeutic Compositions and Administration of Large Effective Size Conjugates

The conjugate of the invention is useful for treating the disease indications that are treated with the parent intact antibody. For example, a conjugate derived from an anti-IL-8 antibody or fragment is useful in the treatment of inflammatory disorders as described in Section (II)(5)(B) below. Therapeutic formulations of the conjugate of the invention can be prepared by utilizing the same procedures described for the formulation of the anti-IL-8 antibodies and fragments of the invention in Section (II)(5)(B) below. The conjugate of the invention can be administered in place of the parent antibody for a given disease indication

by modifying the formulation, dosage, administration protocol, and other aspects of a therapeutic regimen as required by the different pharmacodynamic characteristics of the conjugate and as dictated by common medical knowledge and practice.

#### e. Reagent Uses for Large Effective Size Conjugates

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The conjugate of the invention also finds application as a reagent in an animal model system for in vivo study of the biological functions of the antigen recognized by the conjugate. The conjugate would enable the practitioner to inactivate or detect the cognate antigen in circulation or in tissue for a far greater period of time than would be possible with art-known constructs while removing any Fc interaction (which could attend the use of an intact antibody) from the system. In addition, the increased half-life of the conjugate of the invention can be applied advantageously to the induction of tolerance for the underivatized antibody fragment in a test animal by employing the Wie et al., Int. Archs. Allergy Appl. Immunol., 64: 84-99 (1981) method for allergen tolerization, which would permit the practitioner to repeatedly challenge the tolerized animal with the underivatized parental antibody fragment without generating an immune response against the parental fragment.

## 2. HUMANIZED 6G4.2.5 MONOCLONAL ANTIBODIES AND ANTIBODY FRAGMENTS

In one embodiment, the invention provides an antibody fragment or full length antibody comprising a heavy chain comprising the amino acid sequence of amino acids 1-230 (herein referred to as "6G4.2.5HV11") of the humanized anti-IL-8 6G4.2.5v11 heavy chain polypeptide amino acid sequence of Figs. 37A-37B (SEQ ID NO: 75).

The invention encompasses a single chain antibody fragment comprising the 6G4.2.5HV11, with or without any additional amino acid sequence. In one embodiment, the invention provides a single chain antibody fragment comprising the 6G4.2.5HV11 without any associated light chain amino acid sequence, i.e. a single chain species that makes up one half of a Fab fragment.

Further provided herein are an antibody or antibody fragment comprising the 6G4.2.5HV11, and further comprising a light chain comprising the amino acid sequence of amino acids 1-219 (herein referred to as "6G4.2.5LV11") of the humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65).

In one embodiment, the invention provides a single chain antibody fragment wherein the 6G4.2.5HV11 and the 6G4.2.5LV11 are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment comprises the 6G4.2.5HV11 joined to the 6G4.2.5LV11 by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In another embodiment, the single chain antibody fragment is a species comprising the 6G4.2.5HV11 joined to the 6G4.2.5LV11 by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In yet another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the 6G4.2.5HV11 and a second polypeptide

chain comprises the 6G4.2.5LV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the foregoing two-chain antibody fragment is selected from the group consisting of Fab, Fab', Fab'-SH, and F(ab') 2.

The invention also provides an antibody or antibody fragment comprising a heavy chain containing the 6G4.2.5HV11 and optionally further comprising a light chain containing the 6G4.2.5LV11, wherein the heavy chain, and optionally the light chain, is (are) fused to an additional moiety, such as additional immunoglobulin constant domain sequence. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al. (supra).

In a preferred embodiment, the antibody or antibody fragment comprises the 6G4.2.5HV11 in a heavy chain that is fused to or contains a leucine zipper sequence. The leucine zipper can increase the affinity and/or production efficiency of the antibody or antibody fragment of interest. Suitable leucine zipper sequences include the jun and fos leucine zippers taught by Kostelney et al., J. Immunol., 148: 1547-1553 (1992) and the GCN4 leucine zipper described in the Examples below. In a preferred embodiment, the antibody or antibody fragment comprises the 6G4.2.5HV11 fused at its C-terminus to the GCN4 leucine zipper to yield the amino acid sequence of amino acids 1-275 (herein referred to as "6G4.2.5HV11GCN4") of the heavy chain polypeptide amino acid sequence of Figs. 37A-37B (SEQ ID NO: 75).

# 3. <u>VARIANTS OF HUMANIZED 6G4.2.5 MONOCLONAL ANTIBODIES AND ANTIBODY</u> FRAGMENTS

The invention additionally encompasses humanized anti-IL-8 monoclonal antibody and antibody fragments comprising variants of the 6G4.2.5 complementarity determining regions (CDRs) or variants of the 6G4.2.5v11 variable domains which exhibit higher affinity for human IL-8 and/or possess properties that yield greater efficiency in recombinant production processes.

### A. <u>6G4.2.5LV VARIANTS</u>

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In one aspect, the invention provides humanized anti-IL-8 monoclonal antibodies and antibody fragments comprising the complementarity determining regions (referred to herein as the "CDRs of 6G4.2.5LV") L1, L2, and L3 of the 6G4.2.5 light chain variable domain amino acid sequence of Fig. 24, wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48).

In addition, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a humanized light chain variable domain comprising a variant (hereinafter referred to a "6G4.2.5LV CDRs variant") of the complementarity determining regions L1, L2, and L3 of the 6G4.2.5 variable light chain domain amino acid sequence of Fig. 24 (SEQ ID NO: 48). In one embodiment, the

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invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1N35X35") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than Asn (denoted as "X35") is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48). In a preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/LIN35A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48). In another preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1N35E") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Glu is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48).

In a second aspect, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1S26X<sub>26</sub>") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than Ser (denoted as "X<sub>26</sub>") is substituted for Ser at amino acid position 26, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48). In a preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1S26A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for Ser at amino acid position 26, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48).

In a third aspect, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L3H98X<sub>98</sub>") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than His (denoted as "X<sub>98</sub>") is substituted for His at amino acid position 98. In a

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preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody r antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L3H98A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for His at amino acid position 98.

In a fourth aspect, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1S26X<sub>26</sub>,N35X<sub>35</sub>") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than Ser (denoted as "X<sub>26</sub>") is substituted for Ser at amino acid position 26 and any amino acid other than Asn (denoted as "X<sub>35</sub>") is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48). In a preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1S26A,N35A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for Ser at amino acid position 26 and Ala is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48).

In a fifth aspect, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1N35X<sub>35</sub>/L3H98X<sub>98</sub>") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than Asn (denoted as "X<sub>35</sub>") is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than His (denoted as "X<sub>98</sub>") is substituted for His at amino acid position 98. In a preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1N35A/L3H98A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for His at amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for His at amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for His at amino acid position 98.

In a sixth aspect, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1S26X26/L3H98X98")

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wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than Ser (denoted as "X<sub>26</sub>") is substituted for Ser at amino acid position 26, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than His (denoted as "X<sub>98</sub>") is substituted for His at amino acid position 98. In a preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (herein referred to as "6G4.2.5LV/L1S26A/L3H98A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for Ser at amino acid position 26, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for His at amino acid position 98.

In a seventh aspect, the invention provides a variant 6G4.2.5 humanized antibody or antibody 6G4.2.5LV **CDRs** fragment comprising variant (here referred "6G4.2.5LV/L1S26X263N35X35/L3H98X98") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than Ser (denoted as "X<sub>26</sub>") is substituted for Ser at amino acid position 26 and any amino acid other than Asn (denoted as "X<sub>35</sub>") is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that any amino acid other than His (denoted as "X<sub>QR</sub>") is substituted for His at amino acid position 98. In a preferred embodiment, the invention provides a variant 6G4.2.5 humanized antibody or antibody fragment comprising a 6G4.2.5LV CDRs variant (here referred to as "6G4.2.5LV/L1S26A,N35A/L3H98A") wherein L1 corresponds to amino acids 24-39 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for Ser at amino acid position 26 and Ala is substituted for Asn at amino acid position 35, L2 corresponds to amino acids 55-61 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48), and L3 corresponds to amino acids 94-102 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48) with the proviso that Ala is substituted for His at amino acid position 98.

The humanized light chain variable domains of the invention can be constructed by using any of the techniques for antibody humanization known in the art. Humanization can be essentially performed following the method of Winter and co-workers (Jones et al., Nature 321:522 (1986); Riechmann et al., Nature 332:323 (1988); Verhoeyen et al., Science 239:1534 (1988)), by substituting the CDRs of 6G4.2.5LV or the CDRs of a 6G4.2.5LV CDRs variant for the corresponding sequences of a human antibody light chain variable domain. Accordingly, such "humanized" derivatives containing the CDRs of 6G4.2.5LV or the CDRs of a 6G4.2.5VL CDRs variant are chimeric (Cabilly et al., supra). The humanized

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light chain variable domain comprising the CDRs of 6G4.2.5LV or the CDRs of a 6G4.2.5LV CDRs variant can also contain some FR residues that are substituted by residues from analogous sites in the murine 6G4.2.5 antibody light chain variable domain ("6G4.2.5LV"). The complete amino acid sequence of 6G4.2.5LV is set out as amino acids 1-114 of the amino acid sequence of Fig. 24 (SEQ ID NO: 48).

The invention further provides a humanized antibody or antibody fragment comprising a humanized light chain variable domain comprising the CDRs of 6G4.2.5LV or the CDRs of a 6G4.2.5LV CDRs variant as described above, and further comprising a humanized heavy chain variable domain comprising the complementarity determining regions (CDRs) H1, H2, and H3 of the 6G4.2.5 (murine monoclonal antibody) variable heavy chain domain amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50). The above-described H1, H2, and H3 CDRs of the 6G4.2.5 heavy chain variable domain ("6G4.2.5HV") are collectively referred to as the "CDRs of 6G4.2.5HV".

In another embodiment, the invention provides a humanized antibody or antibody fragment comprising a humanized light chain variable domain comprising the CDRs of 6G4.2.5LV or the CDRs of a 6G4.2.5LV CDRs variant as described above, and further comprising a humanized heavy chain variable domain comprising a variant (herein referred to as a "6G4.2.5HV CDRs variant") of the H1, H2, and H3 CDRs of the 6G4.2.5 (murine monoclonal antibody) variable heavy chain domain amino acid sequence of Fig. 25 (SEQ ID NO: 50). In one 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31Z<sub>31</sub>"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50). In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50).

In a second 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54Z<sub>54</sub>"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50). In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the

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amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50).

In a third 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3D100E"), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100.

In a fourth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3R102K"), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102.

In a fifth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3D106E"), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 106.

In a seventh 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3D100E,R102K"), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino acid position 102.

In an eighth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3R102K,D106E"), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

In a ninth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3D100E,D106E"), wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Glu is substituted for Asp at amino acid position 106.

In a tenth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H3D100E,R102K,D106E"),

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wherein H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), wherein H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and wherein H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102, and Glu is substituted for Asp at amino acid position 106.

**CDRs** variant (referred to herein as eleventh 6G4.2.5HV "6G4.2.5HV/H1S31Z31/H2S54Z54"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amin acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50). In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H2S54A"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50).

In a twelfth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H3D100E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100.

In a thirteenth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31Z<sub>31</sub>/H3R102K"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H3R102K"), H1

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correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102.

A fourteenth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31Z<sub>31</sub>/H3D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H3D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 106.

6G4.2.5HV **CDRs** variant (referred to herein as fifteenth "6G4.2.5HV/H1S31Z31/H3D100E,R102K"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z31") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino acid position 102. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H3D100E,R102K"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino acid position 102.

In a sixteenth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31Z<sub>31</sub>/H3R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position

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102 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H3R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

variant (referred herein 6G4.2.5HV **CDRs** seventeenth In "6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid positi n 100 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H3D100E,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Glu is substituted for Asp at amino acid position 106.

(referred herein as **CDRs** variant 6G4.2.5HV eighteenth In "6G4.2.5HV/H1S31Z31/H3D100E,R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid (referred variant 6G4.2.5HV **CDRs** preferred In 106. position "6G4.2.5HV/H1S31A/H3D100E,R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

In a nineteenth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E"), H1 corresponds to amin acids 26-35 of the amino acid sequence of

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Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as " $Z_{54}$ ") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3D100E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100.

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In a twentieth 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54Z<sub>54</sub>/H3R102K"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3R102K"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102.

In a twenty-first 6G4.2.5HV **CDRs** variant (referred herein as to "6G4.2.5HV/H2S54Z<sub>54</sub>/H3D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 106. In a preferred '6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 106.

In a twenty-second 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,R102K"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is

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substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino acid position 102. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3D100E,R102K"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino acid position 102.

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**CDRs** variant (referred to herein as 6G4.2.5HV In twenty-third "6G4.2.5HV/H2S54Z54/H3R102K,D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3R102K,D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

herein (referred twenty-fourth 6G4.2.5HV **CDRs** variant In "6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3D100E,D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Glu is substituted for Asp at amino acid position 106.

In a twenty-fifth 6G4.2.5HV CDRs variant (referred to herein as

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"6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,R102K,D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H2S54A/H3D100E,R102K,D106E"), H1 corresponds to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50), H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

**CDRs** ln twenty-sixth 6G4.2.5HV variant (referred herein "6G4.2.5HV/H1S31Z31/H2S54Z54/H3D100E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z31") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino In a preferred 6G4.2.5HV CDRs variant (referred to herein as 100. "6G4.2.5HV/H1S31A/H2S54A/H3D100E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100.

In a twenty-seventh 6G4.2.5HV CDRs variant (referred to herein as " $6G4.2.5HV/H1S31Z_{31}/H2S54Z_{54}/H3R102K$ "), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as " $Z_{31}$ ") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as " $Z_{54}$ ") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino

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acid position 102. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H2S54A/H3R102K"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102.

(referred herein 6G4.2.5HV variant twenty-eighth **CDRs** In "6G4.2.5HV/H1S31Z31/H2S54Z54/H3D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino In a preferred 6G4.2.5HV CDRs variant (referred to herein as 106. "6G4.2.5HV/H1S31A/H2S54A/H3D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 106.

herein (referred variant twenty-ninth 6G4.2.5HV **CDRs** In "6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,R102K"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino acid position 102. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Lys is substituted for Arg at amino

acid position 102.

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(referred thirtieth 6G4.2.5HV **CDRs** variant herein ln a "6G4.2.5HV/H1S31Z31/H2S54Z54/H3R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z31") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H2S54A/H3R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

(referred herein as 6G4.2.5HV **CDRs** variant ln thirty-first "6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H2S54A/H3D100E,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100 and Glu is substituted for Asp at amino acid position 106.

In a thirty-second 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser

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(denoted as "Z<sub>31</sub>") is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that any amino acid other than Ser (denoted as "Z<sub>54</sub>") is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106. In a preferred 6G4.2.5HV CDRs variant (referred to herein as "6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K,D106E"), H1 correspond to amino acids 26-35 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 31, H2 corresponds to amino acids 50-66 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Ala is substituted for Ser at amino acid position 54, and H3 corresponds to amino acids 99-111 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50) with the proviso that Glu is substituted for Asp at amino acid position 100, Lys is substituted for Arg at amino acid position 102 and Glu is substituted for Asp at amino acid position 106.

As in the humanization of the light chain variable domain described above, a humanized heavy chain variable domain is constructed by substituting the CDRs of 6G4.2.5HV or the CDRs of a 6G4.2.5HV CDRs variant for the corresponding sequences in a human heavy chain variable domain. The humanized heavy chain variable domain comprising the CDRs of 6G4.2.5HV or the CDRs of a 6G4.2.5HV CDRs variant can also contain some FR residues that are substituted by residues from analogous sites in the murine 6G4.2.5 antibody heavy chain variable domain. The complete amino acid sequence of 6G4.2.5HV is set out as amino acids 1-122 of the amino acid sequence of Fig. 25 (SEQ ID NO: 50).

The choice of human variable domains, both light and heavy, to be used in making the humanized antibodies and antibody fragments is very important to reduce antigenicity. According to the so-called "best-fit" method, the sequence of the variable domain of a rodent antibody is screened against the entire library of known human variable-domain sequences. The human sequence which is closest to that of the rodent is then accepted as the human framework (FR) for the humanized antibody (Sims et al., J. Immunol. 151: 2296 (1993); Chothia and Lesk, J. Mol. Biol. 196:901 (1987)). Another method uses a particular framework derived from the consensus sequence of all human antibodies of a particular subgroup of light or heavy chains. The same framework can be used for several different humanized antibodies (Carter et al., Proc. Natl. Acad. Sci. U.S.A. 89:4285 (1992); Presta et al., J. Immunol. 151:2623 (1993)).

It is also important that the antibodies and antibody fragments of the invention be humanized with retention of high affinity for human IL-8 and other favorable biological properties. To achieve this goal, according to a preferred method, the humanized antibodies and antibody fragments of the invention are prepared by a process of analysis of the parental sequences and various conceptual humanized products using three-dimensional models of the parental and humanized sequences. Three-dimensional immunoglobulin models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of the likely

role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influenc the ability of the candidate immunoglobulin to bind its antigen. In this way, FR residues can be selected and combined from the consensus and parental sequences so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved.

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV are collectively referred to herein as "hu6G4.2.5LV".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1N35X<sub>35</sub> are collectively referred to herein as "hu 6G4.2.5LV/L1N35X<sub>35</sub>".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1N35A are collectively referred to herein as "hu6G4.2.5LV/L1N35A".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1N35E are collectively referred to herein as "hu6G4.2.5LV/L1N35E".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1S26X<sub>26</sub> are collectively referred to herein as "hu6G4.2.5LV/L1S26X<sub>26</sub>".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1S26A are collectively referred to herein as "hu6G4.2.5LV/L1S26A".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L3H98X<sub>98</sub> are collectively referred to herein as "hu6G4.2.5LV/L3H98X<sub>98</sub>".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L3H98A are collectively referred to herein as "hu6G4.2.5LV/L3H98A".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of  $6G4.2.5LV/L1S26X_{26},N35X_{35}$  are collectively referred to herein as "hu6G4.2.5LV/L1S26X<sub>26</sub>,N35X<sub>35</sub>".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1S26A,N35A are collectively referred to herein as "hu6G4.2.5LV/L1S26A,N35A".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1N35X<sub>35</sub>/L3H98X<sub>98</sub> are collectively referred to herein as

"hu6G4.2.5LV/L1N35X35/L3H98X98".

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Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1N35A/L3H98A are collectively referred to herein as "hu6G4.2.5LV/L1N35A/L3H98A".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of  $6G4.2.5LV/L1S26X_{26}/L3H98X_{98}$  are collectively referred to herein as "hu $6G4.2.5LV/L1S26X_{26}/L3H98X_{98}$ ".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1S26A/L3H98A are collectively referred to herein as "hu6G4.2.5LV/L1S26A/L3H98A".

Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of  $6G4.2.5LV/L1S26X_{26}$ ,  $N35X_{35}/L3H98X_{98}$  are collectively referred to herein as "hu6 $G4.2.5LV/L1S26X_{26}$ ,  $N35X_{35}/L3H98X_{98}$ ".

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Any and all humanized light chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5LV/L1S26A,N35A/L3H98A are collectively referred to herein as "hu6G4.2.5LV/L1S26A,N35A/L3H98A".

The humanized light chain variable domain amino acid sequences of hu6G4.2.5LV/L1N35X $_{35}$ , hu6G4.2.5LV/L1S26X $_{26}$ , hu6G4.2.5LV/L1S26X $_{26}$ /L3H98X $_{98}$ , hu6G4.2.5LV/L1S26X $_{26}$ ,N35X $_{35}$ /L3H98X $_{98}$ , hu6G4.2.5LV/L1S26X $_{26}$ /L3H98X $_{98}$ , and hu6G4.2.5LV/L1S26X $_{26}$ /N35X $_{35}$ /L3H98X $_{98}$  are collectively referred to herein as "hu6G4.2.5LV/vL1-3X".

The humanized light chain variable domain amino acid sequences of hu6G4.2.5LV/L1N35A, hu6G4.2.5LV/L1S26A, hu6G4.2.5LV/L1S26A/L3H98A, hu6G4.2.5LV/L1S26A,N35A, hu6G4.2.5LV/L1N35A/L3H98A, hu6G4.2.5LV/L1S26A/L3H98A, hu6G4.2.5LV/L1S26A,N35A/L3H98A are collectively referred to herein as "hu6G4.2.5LV/vL1-3A".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV are collectively referred to herein as "hu6G4.2.5HV".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub> are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A are collectively referred to herein as "hu6G4.2.5HV/H1S31A".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub> are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A are collectively referred to herein as "hu6G4.2.5HV/H2S54A".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H3D100E are collectively referred to herein as "hu6G4.2.5HV/H3D100E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H3R102K are collectively referred to herein as "hu6G4.2.5HV/H3R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H3D106E are collectively referred to herein as "hu6G4.2.5HV/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the

CDRs of 6G4.2.5HV/H3D100E,R102K are collectively referred to herein as "hu6G4.2.5HV/H3D100E,R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H3R102K,D106E are collectively referred to herein as

5 "hu6G4.2.5HV/H3R102K,D106E".

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Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H3D100E,D106E are collectively referred to herein as "hu6G4.2.5HV/H3D100E,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H3D100E,R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of  $6G4.2.5HV/H1S31Z_{31}/H2S54Z_{54}$  are collectively referred to herein as

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z31/H3R102K are collectively referred to herein as

20 "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H3R102K".

"hu6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,R102K".

"hu6G4.2.5HV/H1S31Z31/H3R102K,D106E".

"hu6G4.2.5HV/H1S31Z31/H2S54Z54".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H3D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,R102K are collectively referred to herein as

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H3R102K,D106E are collectively referred to herein as

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,D106E".

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Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H3D100E,R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub>/H3R102K are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub>/H3D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub>/H3R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3R102K are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D106E are collectively referred to herein as

30 "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,R102K are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3R102K,D106E".

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Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of  $6G4.2.5HV/H1S31Z_{31}/H2S54Z_{54}/H3D100E.R102K,D106E$  are collectively referred to herein as "hu $6G4.2.5HV/H1S31Z_{31}/H2S54Z_{54}/H3D100E.R102K.D106E$ ".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H3D100E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3D100E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H3R102K are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H3D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H3D100E,R102K are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3D100E,R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H3R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H3D100E,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3D100E,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the

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CDRs of 6G4.2.5HV/H1S31A/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H3D100E,R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A/H3D100E are collectively referred to herein as

5 "hu6G4.2.5HV/H2S54A/H3D100E".

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Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A/H3R102K are collectively referred to herein as "hu6G4.2.5HV/H2S54A/H3R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A/H3D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54A/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A/H3R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54A/H3R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A/H3D100E,D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54A/H3D100E,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H2S54A/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H2S54A/H3D100E,R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3D100E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A/H3D100E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3R102K are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A/H3R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A/H3D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A/H3R102K,D106E".

Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3D100E,D106E are collectively referred to herein as

"hu6G4.2.5HV/H1S31A/H2S54A/H3D100E,D106E".

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Any and all humanized heavy chain variable domain amino acid sequences which comprise the CDRs of 6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K,D106E".

The humanized heavy chain variable domain amino acid sequences of  $hu6G4.2.5HV/H1S31Z_{31}, \ hu6G4.2.5HV/H2S54Z_{54}, \ hu6G4.2.5HV/H3D100E, \ hu6G4.2.5HV/H3R102K, \ hu6G4.2.5HV/H3D100E, \ hu6G4.2.5HV/H1S31Z_{31}/H2S54Z_{54}, \ hu6G4.2.5HV/H1S31Z_{31}/H3D100E, \ hu6G4.2.5HV/H3D100E, \ hu6G$ 

15 <sub>54</sub>/H3D100E,D106E, hu6G4.2.5HV/H2S54Z<sub>54</sub>/H3D100E,R102K,D106E, hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E, hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3R102K, hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D106E, hu6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>/H3D100E,R102K,

 $hu6G4.2.5HV/H1S31Z_{31}/H2S54Z_{54}/H3D100E, D106E, and hu6G4.2.5HV/H1S31Z_{31}/H2S54Z_{31}/H2S5Z_{31}/H2S5Z_{31}/H2S5Z_{31}/H2S5Z_{31}/H2S_{31}/H2S$ 

20 54/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/vH1-3Z".

hu6G4.2.5HV/H1S31Z31/H2S54Z54/H3R102K,D106E,

The humanized heavy chain variable domain amino acid sequences of hu6G4.2.5HV/H1S31A, hu6G4.2.5HV/H2S54A, hu6G4.2.5HV/H3D100E, hu6G4.2.

- 25 hu6G4.2.5HV/H1S31A/H2S54A, hu6G4.2.5HV/H1S31A/H3D100E, hu6G4.2.5HV/H1S31A/H3R102K. hu6G4.2.5HV/H1S31A/H3D106E, hu6G4.2.5HV/H1S31A/H3D100E,R102K, hu6G4.2.5HV/H1S31A/H3R102K,D106E, hu6G4.2.5HV/H1S31A/H3D100E,D106E, hu6G4.2.5HV/H1S31A/H3D100E,R102K,D106E, hu6G4.2.5HV/H2S54A/H3D100E, hu6G4.2.5HV/H2S54A/H3R102K, hu6G4.2.5HV/H2S54A/H3D106E,
- 30 hu6G4.2.5HV/H2S54A/H3R102K,D106E, hu6G4.2.5HV/H2S54A/H3D100E,D106E, hu6G4.2.5HV/H2S54A/H3D100E,R102K,D106E, hu6G4.2.5HV/H1S31A/H2S54A/H3D100E, hu6G4.2.5HV/H1S31A/H2S54A/H3D106E,

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hu6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K, hu6G4.2.5HV/H1S31A/H2S54A/H3R102K,D106E, hu6G4.2.5HV/H1S31A/H2S54A/H3D100E,D106E, and hu6G4.2.5HV/H1S31A/H2S54A/H3D100E,R102K,D106E are collectively referred to herein as "hu6G4.2.5HV/vH1-3A".

The invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/vL1-3X. In another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/vL1-3A. In yet another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35X35. In still another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35A. In a further embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35E.

The invention additionally provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/vL1-3X, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z. In another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5HV/vL1-3A, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV/vH1-3Z. In yet another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/vL1-3A, and further comprises a heavy chain variable domain comprising the hu6G4.2.5LV/vL1-3A, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV/vH1-3A.

In a further embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35X<sub>35</sub>, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z. In another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/N35X<sub>35</sub>, and further comprises a heavy chain variable domain comprises a light chain variable domain comprises a light chain variable domain comprises a light chain variable domain comprises a humanized heavy chain comprising the hu6G4.2.5LV/L1N35X<sub>35</sub> and further comprises a humanized heavy chain comprising the amino acid sequence of 6G4.2.5HV11.

In an additional embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35A, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z. In another embodiment, the invention provides a humanized antibody or antibody fragment that comprises a light chain variable domain comprising the hu6G4.2.5LV/N35A, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV/vH1-3A. In still another embodiment, the humanized antibody or antibody

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fragment comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35A, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV. In a further embodiment, the humanized antibody or antibody fragment comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35E, and further comprises a heavy chain variable domain comprising the hu6G4.2.5HV. In a preferred embodiment, the antibody or antibody fragment comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35A and further comprises a humanized heavy chain comprising the amino acid sequence of 6G4.2.5HV11. In another preferred embodiment, the antibody or antibody fragment comprises a light chain variable domain comprising the hu6G4.2.5LV/L1N35E and further comprises a humanized heavy chain comprising the amino acid sequence of 6G4.2.5HV11.

The invention encompasses a single chain antibody fragment comprising the hu6G4.2.5LV/vL1-3X, with or without any additional amino acid sequence. In one embodiment, the invention provides a single chain antibody fragment comprising the hu6G4.2.5LV/vL1-3X without any associated heavy chain variable domain amino acid sequence, i.e. a single chain species that makes up one half of an Fv fragment. In another embodiment, the invention provides a single chain antibody fragment comprising the hu6G4.2.5LV/vL1-3A without any associated heavy chain variable domain amino acid sequence. In still another embodiment, the invention provides a single chain antibody fragment comprising the hu6G4.2.5LV/L1N35X<sub>35</sub> without any associated heavy chain variable domain amino acid sequence. In a preferred embodiment, the invention provides a single chain antibody fragment comprising the hu6G4.2.5LV/L1N35A without any associated heavy chain variable domain amino acid sequence. In another preferred embodiment, the invention provides a single chain antibody fragment comprising the hu6G4.2.5LV/L1N35E without any associated heavy chain variable domain amino acid sequence.

In one embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/vL1-3X and the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/vL1-3X joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/vL1-3X joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In another embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/vL1-3A and the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/vL1-3A joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single

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chain antibody fragment is a species comprising the hu6G4.2.5LV/vL1-3A joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In yet another embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/vL1-3A and the hu6G4.2.5HV/vH1-3A are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/vL1-3A joined to the hu6G4.2.5HV/vH1-3A by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/vL1-3A joined to the hu6G4.2.5HV/vH1-3A by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In still another embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/L1N35X<sub>35</sub> and the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/L1N35X<sub>35</sub> joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/L1N35X<sub>35</sub> joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In a further embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/L1N35X<sub>35</sub> and the hu6G4.2.5HV/vH1-3A are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/L1N35X<sub>35</sub> joined to the hu6G4.2.5HV/vH1-3A by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/L1N35X<sub>35</sub> joined to the hu6G4.2.5HV/vH1-3A by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In an additional embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/L1N35A and the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species

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comprising the hu6G4.2.5LV/L1N35A joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/L1N35A joined to the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

Also provided herein is a single chain antibody fragment wherein the hu6G4.2.5LV/L1N35E and the hu6G4.2.5HV are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/L1N35E joined to the hu6G4.2.5HV by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/L1N35E joined to the hu6G4.2.5HV by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In still another embodiment, the invention provides a single chain antibody fragment wherein the hu6G4.2.5LV/L1N35A and the hu6G4.2.5HV/vH1-3A are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5LV/L1N35A joined to the hu6G4.2.5HV/vH1-3A by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5LV/L1N35A joined to the hu6G4.2.5HV/vH1-3A by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In yet another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/vL1-3X and a second polypeptide chain comprises the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

In still another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/vL1-3X and a second polypeptide chain comprises the hu6G4.2.5HV/vH1-3A and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/vL1-3X and a second polypeptide chain comprises the amino acid sequence of 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

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In a further embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5HV/vL1-3A and a second polypeptide chain comprises the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

In still another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/vL1-3A and a second polypeptide chain comprises the hu6G4.2.5HV/vH1-3A and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/vL1-3A and a second polypeptide chain comprises the amino acid sequence of 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

The invention also encompasses an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35X35 and a second polypeptide chain comprises the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

In still another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35X<sub>35</sub> and a second polypeptide chain comprises the hu6G4.2.5HV/vH1-3A and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35X<sub>35</sub> and a second polypeptide chain comprises the amino acid sequence of 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

The invention further encompasses an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35A and a second polypeptide chain comprises the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

The invention also encompasses an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35E and a second polypeptide chain comprises the hu6G4.2.5HV and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

In still another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35A and a second p lypeptide chain comprises the hu6G4.2.5HV/vH1-3A and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the invention provides an

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antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35A and a second polypeptide chain comprises the amino acid sequence of 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In another preferred embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5LV/L1N35E and a second polypeptide chain comprises the amino acid sequence of 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds.

In a preferred embodiment, any of the foregoing two-chain antibody fragments are selected from the group consisting of Fab, Fab', Fab'-SH, Fv, and F(ab') 2. In another preferred embodiment, the antibody fragment is selected from the group consisting of Fab, Fab', Fab'-SH, Fv, and F(ab') 2, wherein the antibody fragment comprises one polypeptide chain comprising the hu6G4.2.5LV/L1N35X35 and a second polypeptide chain comprising the hu6G4.2.5HV. In yet another preferred embodiment, the antibody fragment is selected from the group consisting of Fab, Fab', Fab'-SH, Fv, and F(ab')2, wherein the antibody fragment comprises one polypeptide chain comprising the hu6G4.2.5LV/L1N35A and a second polypeptide chain comprising the hu6G4.2.5HV. In a further preferred embodiment, the antibody fragment comprises one polypeptide chain comprising the hu6G4.2.5LV/L1N35E and a second polypeptide chain comprising the hu6G4.2.5HV. In still another preferred embodiment, the antibody fragment is a F(ab')2 that comprises one polypeptide chain comprising the hu6G4.2.5LV/L1N35A and a second polypeptide chain comprising the amino acid sequence of 6G4.2.5HV11. In an additional preferred embodiment, the antibody fragment is a F(ab')2 that comprises one polypeptide chain comprising the hu6G4.2.5LV/L1N35E and a second polypeptide chain comprising the amino acid sequence of 6G4.2.5HV11. In an additional preferred embodiment, the antibody fragment is a F(ab')2 that comprises one polypeptide chain comprising the hu6G4.2.5LV/L1N35E and a second polypeptide chain comprising the amino acid sequence of 6G4.2.5HV11.

The invention also provides an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/vL1-3X and optionally further comprising a heavy chain variable domain containing the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z, wherein the light chain variable domain, and optionally the heavy chain variable domain, is (are) fused to an additional moiety, such as a immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including lgG, lgM, lgA, lgD, and lgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat *et al.* 

The invention additionally provides an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/vL1-3X and optionally further comprising a heavy chain variable domain containing the hu6G4.2.5HV/vH1-3A, wherein the light chain variable domain, and

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optionally the heavy chain variable domain, is (are) fused to an additional moiety, such as a immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

The invention further provides an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/L1N35X<sub>35</sub> and optionally further comprising a heavy chain variable domain containing the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z, wherein the light chain variable domain, and optionally the heavy chain variable domain, is (are) fused to an additional moiety, such as a immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

The invention additionally provides an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/L1N35X<sub>35</sub> and optionally further comprising a heavy chain variable domain containing the hu6G4.2.5HV/vH1-3A, wherein the light chain variable domain, and optionally the heavy chain variable domain, is (are) fused to an additional moiety, such as a immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat *et al.* 

The invention also encompasses an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/L1N35A and optionally further comprising a heavy chain variable domain containing the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z, wherein the light chain variable domain, and optionally the heavy chain variable domain, is (are) fused to an additional moiety, such as a immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

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The invention additionally provides an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/L1N35A and optionally further comprising a heavy chain variable domain containing the hu6G4.2.5HV/vH1-3A, wherein the light chain variable domain, and optionally the heavy chain variable domain, is (are) fused to an additional moiety, such as a immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including lgG, lgM, lgA, lgD, and lgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

The invention additionally encompasses an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/L1N35A and optionally further comprising a heavy chain containing the amino acid sequence of 6G4.2.5HV11, wherein the light chain variable domain, and optionally the heavy chain, is (are) fused to an additional moiety, such as immunoglobulin constant domain sequences. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

The invention further encompasses an antibody or antibody fragment comprising a light chain variable domain containing the hu6G4.2.5LV/L1N35E and optionally further comprising a heavy chain containing the amino acid sequence of 6G4.2.5HV11, wherein the light chain variable domain, and optionally the heavy chain, is (are) fused to an additional moiety, such as immunoglobulin constant domain sequences. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

In a preferred embodiment, the antibody or antibody fragment comprises a light chain variable domain containing the hu6G4.2.5LV/vL1-3X, and further comprises the hu6G4.2.5HV or hu6G4.2.5HV/vH1-3Z in a heavy chain that is fused to or contains a leucine zipper sequence. The leucine zipper can increase the affinity or production efficiency of the antibody or antibody fragment of interest. Suitable leucine zipper sequences include the jun and fos leucine zippers taught by Kostelney *et al.*, J. Immunol., 148: 1547-1553 (1992) and the GCN4 leucine zipper described in the Examples below.

In particular, the invention provides an antibody or antibody fragment comprising a light chain

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comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than Asn (denoted as "X<sub>35</sub>") is substituted for Asn at amino acid position 35 (herein referred to as "6G4.2.5LV11N35X<sub>35</sub>").

In another embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than Ser (denoted as "X<sub>26</sub>") is substituted for Ser at amino acid position 26 (herein referred to as "6G4.2.5LV11S26X<sub>26</sub>").

In yet another embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than His (denoted as "X<sub>98</sub>") is substituted for His at amino acid position 98 (herein referred to as "6G4.2.5LV11H98X<sub>98</sub>").

In still another embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than Ser (denoted as "X<sub>26</sub>") is substituted for Ser at amino acid position 26 and any amino acid other than Asn (denoted as "X<sub>35</sub>") is substituted for Asn at amino acid position 35 (herein referred to as "6G4.2.5LV11S26X<sub>26</sub>/N35X<sub>35</sub>").

In a further embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than Asn (denoted as " $X_{35}$ ") is substituted for Asn at amino acid position 35 and any amino acid other than His (denoted as " $X_{98}$ ") is substituted for His at amino acid position 98 (herein referred to as " $6G4.2.5LV11N35X_{35}/H98X_{98}$ ").

In an additional embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than Ser (denoted as " $X_{26}$ ") is substituted for Ser at amino acid position 26 and any amino acid other than His (denoted as " $X_{98}$ ") is substituted for His at amino acid position 98 (herein referred to as " $6G4.2.5LV11S26X_{26}/H98X_{98}$ ").

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The invention also encompasses an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that any amino acid other than Ser (denoted as " $X_{26}$ ") is substituted for Ser at amino acid position 26, any amino acid other than Asn (denoted as " $X_{35}$ ") is substituted for Asn at amino acid position 35 and any amino acid other than His (denoted as " $X_{98}$ ") is substituted for His at amino acid position 98 (herein referred to as " $G_{4.2.5LV11S26X_{26}}$ ").

Additionally, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence (SEQ ID NO: 71) of Fig. 36 (herein referred to as "6G4.2.5LV11N35A").

Further provided herein is an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence (SEQ ID NO: 71) of Fig. 45 (herein referred to as "6G4.2.5LV11N35E").

In another embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that Ala is substituted for Ser at amino acid position 26 (herein referred to as "6G4.2.5LV11S26A").

In yet another embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that Ala is substituted for His at amino acid position 98 (herein referred to as "6G4.2.5LV11H98A").

In still another embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that Ala is substituted for Ser at amino acid position 26 and Ala is substituted for Asn at amino acid position 35 (herein referred to as "6G4.2.5LV11S26A/N35A").

In a further embodiment, the invention provides an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that Ala is substituted for Ser at amino acid position 26 and Ala is substituted for His at amino acid position 98 (herein referred to as "6G4.2.5LV11S26A/H98A").

The invention also encompasses an antibody or antibody fragment comprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that Ala is substituted for Asn at amino acid position 35 and Ala is substituted for His at amino acid position 98 (herein

referred to as "6G4.2.5LV11N35A/H98A").

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The invention further encompasses an antibody or antibody fragment c mprising a light chain comprising the amino acid sequence of amino acids 1-219 of the variant humanized anti-IL-8 6G4.2.5v11 light chain polypeptide amino acid sequence of Fig. 31B (SEQ ID NO: 65) with the proviso that Ala is substituted for Ser at amino acid position 26, Ala is substituted for Asn at amino acid position 35, and Ala is substituted for His at amino acid position 98 (herein referred to as "6G4.2.5LV11S26A/N35A/H98A").

The invention provides a single chain antibody fragment comprising a variant light chain selected from the group consisting of 6G4.2.5LV11N35X<sub>35</sub>, 6G4.2.5LV11S26X<sub>26</sub>, 6G4.2.5LV11H98X<sub>98</sub>, 6G4.2.5LV11S26X<sub>26</sub>/ N35X<sub>35</sub>, 6G4.2.5LV11N35X<sub>35</sub>/ H98X<sub>98</sub>, 6G4.2.5LV11S26X<sub>26</sub>/H98X<sub>98</sub>, and 6G4.2.5LV11S26X<sub>26</sub>/ N35X<sub>35</sub>/H98X<sub>98</sub>, with or without any additional amino acid sequence. It will be understood that the group consisting of 6G4.2.5LV11N35X<sub>35</sub>, 6G4.2.5LV11S26X<sub>26</sub>, 6G4.2.5LV11H98X 98, 6G4.2.5LV11S26X<sub>26</sub>/ N35X<sub>35</sub>, 6G4.2.5LV11N35X<sub>35</sub>/ H98X<sub>98</sub>, 6G4.2.5LV11S26X<sub>26</sub>/H98X<sub>98</sub>, and 6G4.2.5LV11S26X<sub>26</sub>/ N35X<sub>35</sub>/H98X<sub>98</sub>, is collectively referred to herein as the "group of 6G4.2.5LV11X variants", and that individual members of this group are generically referred to herein as a "6G4.2.5LV11X variant." In one embodiment, the invention provides a single chain antibody fragment comprising a 6G4.2.5LV11X variant without any associated heavy chain amino acid sequence, i.e. a single chain species that makes up one half of a Fab fragment. In a preferred embodiment, the invention provides a 6G4.2.5LV11N35X<sub>35</sub> variant without any associated heavy chain amino acid sequence.

The invention encompasses a single chain antibody fragment comprising a variant light chain selected from the group consisting of 6G4.2.5LV11N35A, 6G4.2.5LV11S26A, 6G4.2.5LV11H98A, 6G4.2.5LV11S26A/H98A, H98A. 6G4.2.5LV11N35A/ N35A, 6G4.2.5LV11S26A/ 6G4.2.5LV11S26A/ N35A/H98A, with or without any additional amino acid sequence. It will be understood that the group consisting of 6G4.2.5LV11N35A, 6G4.2.5LV11S26A, 6G4.2.5LV11H98A, 6G4.2.5LV11S26A/H98A, 6G4.2.5LV11N35A/ H98A, 6G4.2.5LV11S26A/ N35A, 6G4.2.5LV11S26A/ N35A/H98A is collectively referred to herein as the "group of 6G4.2.5LV11A variants", and that individual members of this group are generically referred to herein as a "6G4.2.5LV11A In one embodiment, the invention provides a single chain antibody fragment comprising a 6G4.2.5LV11A variant without any associated heavy chain amino acid sequence, i.e. a single chain species that makes up one half of a Fab fragment. In a preferred embodiment, the invention provides the 6G4.2.5LV11N35A without any associated heavy chain amino acid sequence.

Further provided herein are an antibody or antibody fragment comprising a light chain comprising a 6G4.2.5LV11X variant, and further comprising a heavy chain comprising the 6G4.2.5HV11. In a preferred embodiment, the invention provides an antibody or antibody fragment comprising a 6G4.2.5LV11N35X<sub>35</sub> variant and further comprising the 6G4.2.5HV11. In a preferred embodiment, the

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invention provides an antibody or antibody fragment comprising the 6G4.2.5LV11N35A and further comprising the 6G4.2.5HV11. In another preferred embodiment, the invention provides an antibody or antibody fragment comprising the 6G4.2.5LV11N35E and further comprising the 6G4.2.5HV11.

In one embodiment, the invention provides a single chain antibody fragment wherein a 6G4.2.5LV11X variant and the 6G4.2.5HV11 are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment comprises a 6G4.2.5LV11X variant joined to the 6G4.2.5HV11 by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In another embodiment, the single chain antibody fragment is a species comprising a 6G4.2.5LV11X variant joined to the 6G4.2.5HV11 by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In still another embodiment, the invention provides a single chain antibody fragment wherein a 6G4.2.5LV11N35X<sub>35</sub> variant and the 6G4.2.5HV11 are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment comprises a 6G4.2.5LV11N35X<sub>35</sub> variant joined to the 6G4.2.5HV11 by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In another embodiment, the single chain antibody fragment is a species comprising a 6G4.2.5LV11N35X<sub>35</sub> variant joined to the 6G4.2.5HV11 by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In a further embodiment, the invention provides a single chain antibody fragment wherein the 6G4.2.5LV11N35A and the 6G4.2.5HV11 are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment comprises the 6G4.2.5LV11N35A joined to the 6G4.2.5HV11 by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In another embodiment, the single chain antibody fragment is a species comprising the 6G4.2.5LV11N35A joined to the 6G4.2.5HV11 by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In an additional embodiment, the invention provides a single chain antibody fragment wherein the 6G4.2.5LV11N35E and the 6G4.2.5HV11 are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment comprises the 6G4.2.5LV11N35E joined to the 6G4.2.5HV11 by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In another embodiment, the single chain antibody fragment is a species comprising the 6G4.2.5LV11N35E

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j ined to the 6G4.2.5HV11 by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

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In yet another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises a 6G4.2.5LV11X variant and a second polypeptide chain comprises the 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In still another embodiment, the invention provides an antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises a 6G4.2.5LV11N35X35 variant and a second polypeptide chain comprises the 6G4.2.5HV11 and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, any of the foregoing two-chain antibody fragments is selected from the group consisting of Fab, Fab', Fab'-SH, and F(ab')2. In still another preferred embodiment, the two-chain antibody fragment is a F(ab')<sub>2</sub> wherein one polypeptide chain comprises the 6G4.2.5LV11N35A and the second polypeptide chain comprises the 6G4.2.5HV11. In a further preferred embodiment, the antibody fragment is a Fab, Fab', Fab'-SH, or F(ab')2 wherein one polypeptide chain comprises the 6G4.2.5LV11N35E and the second polypeptide chain comprises the 6G4.2.5HV11. A particularly preferred embodiment, the antibody fragment is the 6G4V11N35A F(ab')2 GCN4 leucine zipper species described in the Examples below. In another particularly preferred embodiment, the antibody fragment is the 6G4V11N35E F(ab')2 GCN4 leucine zipper species described in the Examples below. In yet another particularly preferred embodiment, the antibody fragment is the 6G4V11N35E Fab described in the Examples below.

The invention also provides an antibody or antibody fragment comprising a light chain containing a 6G4.2.5LV11X variant and optionally further comprising a heavy chain containing the 6G4.2.5HV11, wherein the light chain, and optionally the heavy chain, is (are) fused to an additional moiety, such as additional immunoglobulin constant domain sequence. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

The invention additionally provides an antibody or antibody fragment comprising a light chain containing a 6G4.2.5LV11N35X<sub>35</sub> variant and optionally further comprising a heavy chain containing the 6G4.2.5HV11, wherein the light chain, and optionally the heavy chain, is (are) fused to an additional moiety, such as additional immunoglobulin constant domain sequence. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose,

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including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat *et al.* 

The invention further provides an antibody or antibody fragment comprising a light chain containing the 6G4.2.5LV11N35A and optionally further comprising a heavy chain containing the 6G4.2.5HV11, wherein the light chain, and optionally the heavy chain, is (are) fused to an additional moiety, such as additional immunoglobulin constant domain sequence. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

The invention further provides an antibody or antibody fragment comprising a light chain containing the 6G4.2.5LV11N35E and optionally further comprising a heavy chain containing the 6G4.2.5HV11, wherein the light chain, and optionally the heavy chain, is (are) fused to an additional moiety, such as additional immunoglobulin constant domain sequence. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat *et al.* 

In a preferred embodiment, the antibody or antibody fragment comprises a light chain containing a 6G4.2.5LV11X variant, and further comprises the 6G4.2.5HV11 in a heavy chain that is fused to or contains a leucine zipper sequence. The leucine zipper can increase the affinity or production efficiency of the antibody or antibody fragment of interest. Suitable leucine zipper sequences include the jun and fos leucine zippers taught by Kostelney et al., J. Immunol., 148: 1547-1553 (1992) and the GCN4 leucine zipper described in the Examples below. In another preferred embodiment, the antibody or antibody fragment comprises a light chain containing the 6G4.2.5LV11N35A, and further comprises a heavy chain containing the 6G4.2.5HV11 fused to the GCN4 leucine zipper. In yet another preferred embodiment, the antibody or antibody fragment comprises a light chain containing the 6G4.2.5LV11N35E, and further comprises a heavy chain containing the 6G4.2.5HV11 fused to the GCN4 leucine zipper.

## B. <u>6G4.2.5HV VARIANTS</u>

The invention provides humanized antibodies and antibody fragments comprising the CDRs of a 6G4.2.5HV CDR variant. The use of a 6G4.2.5HV CDRs variant in the humanized antibodies and antibody fragments of the invention confer the advantages of higher affinity for human IL-8 and/or improved recombinant manufacturing economy.

A heavy chain variable domain comprising the CDRs of a 6G4.2.5HV CDRs variant can be

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humanized in conjunction with a light chain comprising the CDRs of 6G4.2.5LV or the CDRs of a 6G4.2.5LV CDRs variant, essentially as described in Section (II)(2)(A) above. In one embodiment, the invention provides a humanized antibody or antibody fragment comprising a 6G4.2.5HV CDRs variant selected from the group consisting of 6G4.2.5HV/H1S31Z<sub>31</sub>, 6G4.2.5HV/H2S54Z<sub>54</sub>, and 6G4.2.5HV/H1S31Z<sub>31</sub>/H2S54Z<sub>54</sub>. In addition, the invention provides a humanized antibody or antibody fragment comprising a 6G4.2.5HV CDRs variant selected from the group consisting of 6G4.2.5HV/H1S31A, 6G4.2.5HV/H2S54A, and 6G4.2.5HV/H1S31A/H2S54A. In particular, the 6G4.2.5HV CDRs variants can be used to construct a humanized antibody or antibody comprising the hu6G4.2.5HV/vH1-3Z as described in Section (II)(2)(A) above.

The invention additionally provides a humanized antibody or antibody fragment that comprises a heavy chain variable domain comprising the hu6G4.2.5HV/vH1-3Z, and further comprises a light chain variable domain comprising the hu6G4.2.5LV or hu6G4.2.5LV/vL1-3X.

The invention further encompasses a single chain humanized antibody fragment comprising the hu6G4.2.5HV/vH1-3Z, with or without any additional amino acid sequence. In one embodiment, the invention provides a single chain antibody fragment comprising the hu6G4.2.5HV/vH1-3Z without any associated heavy chain variable domain amino acid sequence, i.e. a single chain species that makes up one half of an Fv fragment.

In one embodiment, the invention provides a single chain humanized antibody fragment wherein the hu6G4.2.5HV/vH1-3Z and the hu6G4.2.5LV or hu6G4.2.5LV/vL1-3X are contained in a single chain polypeptide species. In a preferred embodiment, the single chain antibody fragment is a scFv species comprising the hu6G4.2.5HV/vH1-3Z joined to the hu6G4.2.5LV or hu6G4.2.5LV/vL1-3X by means of a flexible peptide linker sequence, wherein the heavy chain and light chain variable domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fv species. In another embodiment, the single chain antibody fragment is a species comprising the hu6G4.2.5HV/vH1-3Z joined to the hu6G4.2.5LV or hu6G4.2.5LV/vL1-3X by a linker that is too short to permit intramolecular pairing of the two variable domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In yet another embodiment, the invention provides a humanized antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises the hu6G4.2.5HV/vH1-3Z and a second polypeptide chain comprises the hu6G4.2.5LV or hu6G4.2.5LV/vL1-3X and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the foregoing two-chain antibody fragment is selected from the group consisting of Fab, Fab', Fab'-SH, Fv, and F(ab') 2.

The invention also provides a humanized antibody or antibody fragment comprising a heavy chain variable domain containing the hu6G4.2.5HV/vH1-3Z and optionally further comprising a light chain variable domain containing the hu6G4.2.5LV or hu6G4.2.5LV/vL1-3X, wherein the heavy chain variable

domain, and optionally the light chain variable domain, is (are) fused to an additional moiety, such as an immunoglobulin constant domain. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat et al.

In a preferred embodiment, the humanized antibody or antibody fragment comprises the hu6G4.2.5HV/vH1-3Z in a heavy chain that is fused to or contains a leucine zipper sequence. The leucine zipper can increase the affinity or production efficiency of the antibody or antibody fragment of interest. Suitable leucine zipper sequences include the jun and fos leucine zippers taught by Kostelney et al., J. Immunol., 148: 1547-1553 (1992) and the GCN4 leucine zipper described in the Examples below.

In addition, the invention provides a humanized antibody or antibody fragment comprising a heavy chain comprising the amino acid sequence of amino acids 1-230 of the 6G4.2.5HV11 polypeptide amino acid sequence of Figs. 37A-37B (SEQ ID NO: 75) with the proviso that Ala is substituted for Ser at amino acid position 31 (hereinafter referred to as "6G4.2.5HV11S31A").

In another embodiment, the invention provides a humanized antibody or antibody fragment comprising a heavy chain comprising the amino acid sequence of amino acids 1-230 of the 6G4.2.5HV11 polypeptide amino acid sequence of Figs. 37A-37B (SEQ ID NO: 75) with the proviso that Ala is substituted for Ser at amino acid position 54 (hereinafter referred to as "6G4.2.5HV11S54A").

In yet another embodiment, the invention provides a humanized antibody or antibody fragment comprising a heavy chain comprising the amino acid sequence of amino acids 1-230 of the 6G4.2.5HV11 polypeptide amino acid sequence of Figs. 37A-37B (SEQ ID NO: 75) with the proviso that Ala is substituted for Ser at amino acid position 31 and Ala is substituted for Ser at amino acid position 54 (hereinafter referred to as "6G4.2.5HV11S31A/S54A").

Further provided herein is a humanized antibody or antibody fragment that comprises any of the light and heavy chain combinations listed in Tables 1 and 2 below.

Table 1

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	Heavy Chain	Light Chain
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	6G4.2.5HV11S31A	6G4.2.5LV11
	6G4.2.5HV11S31A	6G4.2.5LV11N35A
	. 6G4.2.5HV11S31A	6G4.2.5LV11S26A
	6G4.2.5HV11S31A	6G4.2.5LV11H98A
35	6G4.2.5HV11S31A	6G4.2.5LV11S26A/N35A
	6G4.2.5HV11S31A	6G4.2.5LV11S26A/H98A
	6G4.2.5HV11S31A	6G4.2.5LV11N35A/H98A
	6G4.2.5HV11S31A	6G4.2.5LV11S26A/N35A/H98A
	6G4.2.5HV11S54A	6G4.2.5LV11
40	6G4.2.5HV11S54A	6G4.2.5LV11N35A
	6G4.2.5HV11S54A	6G4.2.5LV11S26A
	6G4.2.5HV11S54A	6G4.2.5LV11H98A

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## Table 2

	Table	2	
	Heavy Chain	Li	ght Chain
	6G4.2.5HV11S54A		G4.2.5LV11S26A/N35A
5	6G4.2.5HV11S54A		54.2.5LV11S26A/H98A
Ū	6G4.2.5HV11S54A		34.2.5LV11N35A/H98A
	6G4.2.5HV11S54A		G4.2.5LVI 1S26A/N35A/H98A
	6G4.2.5HV11S31A/S54A		G4.2.5LV11
	6G4.2.5HV11S31A/S54A		G4.2.5LV11N35A G4.2.5LV11S26A
10	6G4.2.5HV11S31A/S54A		G4.2.5LV11H98A
	6G4.2.5HV11S31A/S54A 6G4.2.5HV11S31A/S54A	<del>-</del>	4.2.5LV11S26A/N35A
	6G4.2.5HV11S31A/S54A	_	G4.2.5LV11S26A/H98A
	6G4.2.5HV11S31A/S54A		G4.2.5LV11N35A/H98A
15	6G4.2.5HV11S31A/S54A		G4.2.5LV11S26A/N35A/H98A
	6G4.2.5HV11S31A	6	G4.2.5LV11
	6G4.2.5HV11S31A	6	G4.2.5LV11N35X <sub>35</sub>
	6G4.2.5HV11S31A	6	G4.2.5LV11S26X <sub>26</sub>
	6G4.2.5HV11S31A		G4.2.5LV11H98X <sub>98</sub>
20	6G4.2.5HV11S31A		G4.2.5LV11S26X <sub>26</sub> /N35X <sub>35</sub>
	6G4.2.5HV11S31A	6	G4.2.5LV11S26X <sub>26</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S31A		G4.2.5LV11N35X <sub>35</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S31A	6	G4.2.5LV11S26X <sub>26</sub> /N35X <sub>35</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S54A	6	5G4.2.5LV11
25	6G4.2.5HV11S54A	6	5G4.2.5LV11N35X <sub>35</sub>
	6G4.2.5HV11S54A	6	5G4.2.5LV11S26X <sub>26</sub>
	6G4.2.5HV11S54A	Ć	6G4.2.5LV11H98X <sub>98</sub>
	6G4.2.5HV11S54A	(	6G4.2.5LV11S26X <sub>26</sub> /N35X <sub>35</sub>
	6G4.2.5HV11S54A	(	6G4.2.5LV11S26X <sub>26</sub> /H98X <sub>98</sub>
30	6G4.2.5HV11S54A		6G4.2.5LV11N35X <sub>35</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S54A	•	6G4.2.5LV11S26X <sub>26</sub> /N35X <sub>35</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S31A/S54A	•	6G4.2.5LV11
	6G4.2.5HV11S31A/S54A		6G4.2.5LV11N35X <sub>35</sub>
	6G4.2.5HV11S31A/S54A	1	6G4.2.5LV11S26X <sub>26</sub>
35	6G4.2.5HV11S31A/S54A		6G4.2.5LV11H98X <sub>98</sub>
	6G4.2.5HV11S31A/S54A		6G4.2.5LV11S26X <sub>26</sub> /N35X <sub>35</sub>
	6G4.2.5HV11S31A/S54A		6G4.2.5LV11S26X <sub>26</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S31A/S54A		6G4.2.5LV11N35X <sub>35</sub> /H98X <sub>98</sub>
	6G4.2.5HV11S31A/S54A		6G4.2.5LV11S26X <sub>26</sub> /N35X <sub>35</sub> /H98X <sub>98</sub>
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The invention encompasses a single chain humanized antibody fragment comprising a variant heavy chain selected from the group consisting of 6G4.2.5HV11S31A, 6G4.2.5HV11S54A, and 6G4.2.5HV11S31A/S54A, with or without any additional amino acid sequence. It will be understood that the group consisting of 6G4.2.5HV11S31A, 6G4.2.5HV11S54A, and 6G4.2.5HV11S31A/S54A is collectively referred to herein as the "group of 6G4.2.5HV11A variants", and that individual members of

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this group are generically referred to herein as a "6G4.2.5HV11A variant." In one embodiment, the invention provides a single chain humanized antibody fragment comprising a 6G4.2.5HV11A variant without any associated light chain amino acid sequence, i.e. a single chain species that makes up one half of a Fab fragment.

Further provided herein are a humanized antibody or antibody fragment comprising a heavy chain comprising a 6G4.2.5HV11A variant, and further comprising a light chain comprising a 6G4.2.5LV11A variant or a 6G4.2.5LV11X variant. In another embodiment, the humanized antibody or antibody fragment comprises any combination of light and heavy chains listed in Tables 1 and 2 above. In one embodiment, the invention provides a humanized antibody or antibody fragment comprising a 6G4.2.5HV11A variant and further comprising the 6G4.2.5LV11N35X<sub>35</sub>. In a preferred embodiment, the invention provides a humanized antibody or antibody fragment comprising a 6G4.2.5HV11A variant and further comprising the 6G4.2.5LV11N35A.

In yet another embodiment, the invention provides a single chain humanized antibody fragment wherein a 6G4.2.5HV11A variant and the 6G4.2.5LV11 are contained in a single chain polypeptide species. In another embodiment, the invention provides a single chain humanized antibody fragment wherein any pair of light and heavy chains listed in Tables 1 and 2 above is contained in a single chain polypeptide species. In yet another embodiment, the invention provides a single chain humanized antibody fragment wherein a 6G4.2.5HV11A variant and a 6G4.2.5LV11X variant are contained in a single chain polypeptide species. In still another embodiment, the invention provides a single chain humanized antibody fragment wherein a 6G4.2.5HV11A variant and a 6G4.2.5LV11N35X35 variant are contained in a single chain polypeptide species. In an additional embodiment, the invention provides a single chain humanized antibody fragment wherein a 6G4.2.5HV11A variant and the 6G4.2.5LV11N35A variant are contained in a single chain polypeptide species.

In a preferred embodiment, the single chain humanized antibody fragment comprises a 6G4.2.5HV11A variant joined to a 6G4.2.5LV11X variant, 6G4.2.5LV11N35X<sub>35</sub> variant, 6G4.2.5LV11N35A variant, or 6G4.2.5LV11 by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In a further embodiment, the single chain humanized antibody fragment is a species comprising a 6G4.2.5HV11A variant joined to a 6G4.2.5LV11X variant, 6G4.2.5LV11N35X<sub>35</sub> variant, 6G4.2.5LV11N35A variant, or 6G4.2.5LV11 by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In still another embodiment, the single chain humanized antibody fragment comprises any pair of light and heavy chains listed in Tables 1 and 2 above joined by means of a flexible peptide linker sequence, wherein the heavy chain and light chain domains can associate in a "dimeric" structure analogous to that formed in a two-chain Fab species. In an additional embodiment, the single chain humanized antibody

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fragment comprises any pair of light and heavy chains listed in Tables 1 and 2 above joined by a linker that is too short to permit intramolecular pairing of complementary domains, i.e. a single chain polypeptide monomer that forms a diabody upon dimerization with another monomer.

In yet another embodiment, the invention provides a humanized antibody fragment comprising a plurality of polypeptide chains, wherein one polypeptide chain comprises a 6G4.2.5HV11A variant and a second polypeptide chain comprises a 6G4.2.5LV11X variant, 6G4.2.5LV11N35X<sub>35</sub> variant, 6G4.2.5LV11N35A variant, or 6G4.2.5LV11, and the two polypeptide chains are covalently linked by one or more interchain disulfide bonds. In a preferred embodiment, the foregoing two-chain antibody fragment is selected from the group consisting of Fab, Fab', Fab'-SH, and F(ab')<sub>2</sub>.

In an additional embodiment, the invention provides a two-chain humanized antibody fragment comprising any pair of heavy and light chains listed in Tables 1 and 2 above, wherein each chain is contained on a separate molecule. In another embodiment, the two-chain antibody fragment comprising any pair of heavy and light chains listed in Tables 1 and 2 above is selected from the group consisting of Fab, Fab', Fab'-SH, and F(ab') 2. In a preferred embodiment, the two-chain humanized antibody fragment is a F(ab') 2 comprising any pair of heavy and light chains listed in Tables 1 and 2 above. In another preferred embodiment, the two-chain humanized antibody fragment is a F(ab') 2 wherein one polypeptide chain comprises a 6G4.2.5HV11A variant and the second polypeptide chain comprises the 6G4.2.5LV11N35A.

The invention also provides a humanized antibody or antibody fragment comprising a heavy chain containing a 6G4.2.5HV11A variant and optionally further comprising a light chain containing a 6G4.2.5LV11X variant, 6G4.2.5LV11N35X<sub>35</sub> variant, 6G4.2.5LV11N35A, or 6G4.2.5HV11, wherein the heavy chain, and optionally the light chain, is (are) fused to an additional moiety, such as additional immunoglobulin constant domain sequence. Constant domain sequence can be added to the heavy chain and/or light chain sequence(s) to form species with full or partial length heavy and/or light chain(s). It will be appreciated that constant regions of any isotype can be used for this purpose, including IgG, IgM, IgA, IgD, and IgE constant regions, and that such constant regions can be obtained from any human or animal species. Preferably, the constant domain sequence is human in origin. Suitable human constant domain sequences can be obtained from Kabat *et al.* (supra).

In a preferred embodiment, the humanized antibody or antibody fragment comprises a 6G4.2.5HV11A variant in a heavy chain that is fused to or contains a leucine zipper sequence. The leucine zipper can increase the affinity or production efficiency of the antibody or antibody fragment of interest. Suitable leucine zipper sequences include the jun and fos leucine zippers taught by Kostelney et al., 1. Immunol., 148: 1547-1553 (1992) and the GCN4 leucine zipper described in the Examples below.

## C. Bispecific Antibodies

Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, ne of the binding

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specifications is for IL-8, the other one is for any other antigen. For example, bispecific antibodies specifically binding a IL-8 and neurotrophic factor, or two different types of IL-8 polypeptides are within the scope of the present invention.

Methods for making bispecific antibodies are known in the art. Traditionally, the recombinant production of bispecific antibodies is based on the co-expression of two immunoglobulin heavy chain-light chain pairs, where the two heavy chains have different specificities (Milstein and Cuello, Nature 305:537 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of 10 different antibody molecules, of which only one has the correct bispecific structure. The purification of the correct molecule, which is usually done by affinity chromatography steps, is rather cumbersome, and the product yields are low. Similar procedures are disclosed in WO 93/08829 published 13 May 1993, and in Traunecker et al., EMBO J. 10:3655 (1991).

According to a different and more preferred approach, antibody-variable domains with the desired binding specificities (antibody-antigen combining sites) are fused to immunoglobulin constant-domain sequences. The fusion preferably is with an immunoglobulin heavy-chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1), containing the site necessary for light-chain binding, present in at least one of the fusions. DNAs encoding the immunoglobulin heavy chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host organism. This provides for great flexibility in adjusting the mutual proportions of the three polypeptide fragments in embodiments when unequal ratios of the three polypeptide chains used in the construction provide the maximum yields. It is, however, possible to insert the coding sequences for two or all three polypeptide chains in one expression vector when the production of at least two polypeptide chains in equal ratios results in high yields or when the ratios are of no particular significance. In a preferred embodiment of this approach, the bispecific antibodies are composed of a hybrid immunoglobulin heavy chain with a first binding specificity in one arm, and a hybrid immunoglobulin heavy chain-light chain pair (providing a second binding specificity) in the other arm. This asymmetric structure facilitates the separation of the desired bispecific compound from unwanted immunoglobulin chain combinations, as the presence of an immunoglobulin light chain in only one half of the bispecific molecule provides for a facile way of separation. For further details of generating bispecific antibodies, see, for example, Suresh et al., Methods in Enzymology 121:210 (1986).

According to another approach, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the C<sub>H</sub>3 domain of an antibody constant domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). This provides a mechanism for increasing the yield of the heterodimer over other unwanted end-products such as

homodimers.

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Bispecific antibodies include cross-linked or "heteroconjugate" antibodies. For example, one of the antibodies in the heteroconjugate can be coupled to avidin, the other to biotin. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (US Patent No. 4,676,980), and for treatment of HIV infection (WO 91/00360, WO 92/00373, and EP 03089). Heteroconjugate antibodies may be made using any convenient cross-linking methods. Suitable cross-linking agents are well known in the art, and are disclosed in US Patent No. 4,676,980, along with a number of cross-linking techniques.

Techniques for generating bispecific antibodies from antibody fragments have also been described in the literature. For example, bispecific antibodies can be prepared using chemical linkage. Brennan et al., Science, 229: 81 (1985) describe a procedure wherein intact antibodies are proteolytically cleaved to generate F(ab') 2 fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

Recent progress has facilitated the direct recovery of Fab'-SH fragments from E. coli, which can be chemically coupled to form bispecific antibodies. Shalaby et al., J. Exp. Med., 175: 217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab') 2 molecule. Each Fab' fragment was separately secreted from E. coli and subjected to directed chemical coupling in vitro to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the HER2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets.

Various techniques for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelny et al., J. Immunol., 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., Proc. Natl. Acad. Sci. USA, 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a heavy-chain variable domain (VH) connected to a light-chain variable domain (VL) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the VH and VL domains of one fragment are forced to pair with the complementary VL and VH domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See Gruber et al., J. Immunol., 152:5368 (1994).

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Antibodies with more than two valencies are contemplated. For example, trispecific antibodies can be prepared. Tutt et al. J. Immunol. 147: 60 (1991).

# 4. <u>Production of Humanized Anti-IL-8 6G4.2.5 Monoclonal Antibody, Antibody Fragments, and Variants</u>

The antibodies and antibody fragments of the invention can be produced using any convenient antibody manufacturing process known in the art. Typically, the antibody or antibody fragment is made using recombinant expression systems. A multiple polypeptide chain antibody or antibody fragment species can be made in a single host cell expression system wherein the host cell produces each chain of the antibody or antibody fragment and assembles the polypeptide chains into a multimeric structure to form the antibody or antibody fragment in vivo, followed by recovery of the antibody or antibody fragment from the host cell. For example, suitable recombinant expression systems for the production of complete antibody or antibody fragment are described in Lucas et al., Nucleic Acids Res., 24: 1774-1779 (1996). Alternatively, the separate polypeptide chains of the desired antibody or antibody fragment can be made in separate expression host cells, separately recovered from the respective host cells, and then mixed in vitro under conditions permitting the formation of the multi-subunit antibody or antibody fragment of interest. For example, U.S. Pat. No. 4,816,567 to Cabilly et al. and Carter et al., Bio/Technology, 10: 163-167 (1992) provide methods for recombinant production of antibody heavy and light chains in separate expression hosts followed by assembly of antibody from separate heavy and light chains in vitro.

The following discussion of recombinant expression methods applies equally to the production of single chain antibody polypeptide species and multi-subunit antibody and antibody fragment species. All recombinant procedures for the production of antibody or antibody fragment provided below shall be understood to describe: (1) manufacture of single chain antibody species as the desired end-product; (2) manufacture of multi-subunit antibody or antibody fragment species by production of all subunits in a single host cell, subunit assembly in the host cell, optionally followed by host cell secretion of the multi-subunit end-product into the culture medium, and recovery of the multi-subunit end-product from the host cell and/or culture medium; and (3) manufacture of multi-subunit antibody or antibody fragment by production of subunits in separate host cells (optionally followed by host cell secretion of subunits into the culture medium), recovery of subunits from the respective host cells and/or culture media, followed by in vitro subunit assembly to form the multi-subunit end-product. In the case of a multi-subunit antibody or antibody fragment produced in a single host cell, it will be appreciated that production of the various subunits can be effected by expression of multiple polypeptide-encoding nucleic acid sequences carried on a single vector or by expression of polypeptide-encoding nucleic acid sequences carried on multiple vectors contained in the host cell.

# A. Construction of DNA Encoding Humanized 6G4.2.5 Monoclonal Antibodies, Antibody Fragments, and Variants

Following the selection of the humanized antibody or antibody fragment of the invention according to the methods described above, the practitioner can use the genetic code to design DNAs

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encoding the desired antibody or antibody fragment. In one embodiment, codons preferred by the expression host cell are used in the design of a DNA encoding the antibody or antibody fragment of interest. DNA encoding the desired antibody or antibody fragment can be prepared by a variety of methods known in the art. These methods include, but are not limited to, chemical synthesis by any of the methods described in Engels et al., Agnew. Chem. Int. Ed. Engl., 28: 716-734 (1989), the entire disclosure of which is incorporated herein by reference, such as the triester, phosphite, phosphoramidite and H-phosphonate methods.

A variation on the above procedures contemplates the use of gene fusions, wherein the gene(s) encoding the antibody or antibody fragment is associated, in the vector, with a gene encoding another protein or a fragment of another protein. This results in the antibody or antibody fragment being produced by the host cell as a fusion with another protein. The "other" protein is often a protein or peptide which can be secreted by the cell, making it possible to isolate and purify the desired protein from the culture medium and eliminating the necessity of destroying the host cells which arises when the desired protein remains inside the cell. Alternatively, the fusion protein can be expressed intracellularly. It is advantageous to use fusion proteins that are highly expressed.

The use of gene fusions, though not essential, can facilitate the expression of heterologous proteins in *E. coli* as well as the subsequent purification of those gene products (Harris, T. J. R. in *Genetic Engineering*, Williamson, R., Ed., Academic, London, Vol. 4, p. 127(1983); Uhlen, M. & Moks, T., *Methods Enzymol.* 185:129-143 (1990)). Protein A fusions are often used because the binding of protein A, or more specifically the Z domain of protein A, to IgG provides an "affinity handle" for the purification of the fused protein (Nilsson, B. & Abrahmsen, L. *Methods Enzymol.* 185:144-161 (1990)). It has also been shown that many heterologous proteins are degraded when expressed directly in *E. coli*, but are stable when expressed as fusion proteins (Marston, F. A. O., *Biochem J.* 240: 1 (1986)).

Fusion proteins can be cleaved using chemicals, such as cyanogen bromide, which cleaves at a methionine, or hydroxylamine, which cleaves between an Asn and Gly. Using standard recombinant DNA methodology, the nucleotide base pairs encoding these amino acids may be inserted just prior to the 5' end of the antibody or antibody fragment gene(s).

Alternatively, one can employ proteolytic cleavage of fusion proteins, which has been recently reviewed (Carter, P. (1990) in *Protein Purification: From Molecular Mechanisms to Large-Scale Processes*. Ladisch, M. R., Willson, R. C., Painton, C. C., and Builder, S. E., eds., American Chemical Society Symposium Series No. 427, Ch 13, 181-193).

Proteases such Factor Xa, thrombin, subtilisin and mutants thereof, have been successfully used to cleave fusion proteins. Typically, a peptide linker that is amenable to cleavage by the protease used is inserted between the "other" protein (e.g., the Z domain of protein A) and the protein of interest, such as humanized anti-IL-8 antibody or antibody fragment. Using recombinant DNA methodology, the nucleotide base pairs encoding the linker are inserted between the genes or gene fragments coding for the other proteins. Proteolytic cleavage of the partially purified fusion protein containing the correct linker can then

be carried out on either the native fusion protein, or the reduced or denatured fusion protein.

Various techniques are also available which may now be employed to produce variant humanized antibodies or antibody fragments, which encodes for additions, deletions, or changes in amino acid sequence of the resultant protein(s) relative to the parent humanized antibody or antibody fragment.

By way of illustration, with expression vectors encoding humanized antibody or antibody fragment in hand, site specific mutagenesis (Kunkel et al., Methods Enzymol. 204:125-139 (1991); Carter, P., et al., Nucl. Acids. Res. 13:4331 (1986); Zoller, M. J. et al., Nucl. Acids Res. 10:6487 (1982)), cassette mutagenesis (Wells, J. A., et al., Gene 34:315 (1985)), restriction selection mutagenesis (Wells, J. A., et al., Philos. Trans, R. Soc. London SerA 317, 415 (1986)) or other known techniques may be performed on the antibody or antibody fragment DNA. The variant DNA can then be used in place of the parent DNA by insertion into the aforementioned expression vectors. Growth of host bacteria containing the expression vectors with the mutant DNA allows the production of variant humanized antibodies or antibody fragments, which can be isolated as described herein.

## B. Insertion of DNA into a Cloning Vehicle

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The DNA encoding the antibody or antibody fragment is inserted into a replicable vector for further cloning (amplification of the DNA) or for expression. Many vectors are available, and selection of the appropriate vector will depend on (1) whether it is to be used for DNA amplification or for DNA expression, (2) the size of the DNA to be inserted into the vector, and (3) the host cell to be transformed with the vector. Each vector contains various components depending on its function (amplification of DNA or expression of DNA) and the host cell for which it is compatible. The vector components generally include, but are not limited to, one or more of the following: a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence.

## (i) Signal Sequence Component

In general, a signal sequence may be a component of the vector, or it may be a part of the antibody or antibody fragment DNA that is inserted into the vector. Preferably, a heterologous signal sequence selected and fused to the antibody or antibody fragment DNA such that the signal sequence in the corresponding fusion protein is recognized, transported and processed (i.e., cleaved by a signal peptidase) in the host cell's protein secretion system. In the case of prokaryotic host cells, the signal sequence is selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. In a preferred embodiment, the STII signal sequence is used as described in the Examples below. For yeast secretion the native signal sequence may be substituted by, e.g., the yeast invertase leader,  $\alpha$  factor leader (including Saccharomyces and Kluyveromyces  $\alpha$ -factor leaders), or acid phosphatase leader, the C. albicans glucoamylase leader, or the signal described in WO 90/13646. In mammalian cell expression, mammalian signal sequences as well as viral secretory leaders, for example, the herpes simplex gD signal, are available.

## (ii) Origin of Replication Component

Both expression and cloning vectors contain a nucleic acid sequence that enables

the vector to replicate in one or more selected host cells. Generally, in cloning vectors this sequence is one that enables the vector to replicate independently of the host chromosomal DNA, and includes origins of replication or autonomously replicating sequences. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2µ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors (the SV40 origin may typically be used only because it contains the early promoter).

Most expression vectors are "shuttle" vectors, i.e. they are capable of replication in at least one class of organisms but can be transfected into another organism for expression. For example, a vector is cloned in *E. coli* and then the same vector is transfected into yeast or mammalian cells for expression even though it is not capable of replicating independently of the host cell chromosome.

DNA may also be amplified by insertion into the host genome. This is readily accomplished using *Bacillus* species as hosts, for example, by including in the vector a DNA sequence that is homologous to a sequence found in *Bacillus* genomic DNA. Transfection of *Bacillus* with this vector results in homologous recombination with the genome and insertion of the antibody or antibody fragment DNA.

## (iii) Selection Gene Component

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Expression and cloning vectors should contain a selection gene, also termed a selectable marker. This gene encodes a protein necessary for the survival or growth of transformed host cells grown in a selective culture medium. Host cells not transformed with the vector containing the selection gene will not survive in the culture medium. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g. ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, e.g. the gene encoding D-alanine racemase for *Bacilli*.

One example of a selection scheme utilizes a drug to arrest growth of a host cell. Those cells that are successfully transformed with a heterologous gene express a protein conferring drug resistance and thus survive the selection regimen. Examples of such dominant selection use the drugs neomycin (Southern et al., J. Molec. Appl. Genet., 1: 327 (1982)), mycophenolic acid (Mulligan et al., Science, 209: 1422 (1980)) or hygromycin (Sugden et al., Mol. Cell. Biol., 5: 410-413 (1985)). The three examples given above employ bacterial genes under eukaryotic control to convey resistance to the appropriate drug (G418 or neomycin (geneticin), xgpt (mycophenolic acid), and hygromycin, respectively.)

Another example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the antibody or antibody fragment nucleic acid, such as dihydrofolate reductase (DHFR) or thymidine kinase. The mammalian cell transformants are placed under selection pressure which only the transformants are uniquely adapted to survive by virtue of having taken up the marker. Selection pressure is imposed by culturing the transformants under conditions in which the concentration of selection agent in the medium is successively changed, thereby leading to amplification of

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both the selection gene and the DNA that encodes the antibody or antibody fragment. Amplification is the process by which genes in greater demand for the production f a protein critical for growth are reiterated in tandem within the chromosomes of successive generations of recombinant cells. Increased quantities of the antibody or antibody fragment are synthesized from the amplified DNA.

For example, cells transformed with the DHFR selection gene are first identified by culturing all of the transformants in a culture medium that contains methotrexate (Mtx), a competitive antagonist of DHFR. An appropriate host cell when wild-type DHFR is employed is the Chinese hamster ovary (CHO) cell line deficient in DHFR activity, prepared and propagated as described by Urlaub and Chasin, <u>Proc. Natl. Acad. Sci. USA, 77</u>: 4216 (1980). The transformed cells are then exposed to increased levels of methotrexate. This leads to the synthesis of multiple copies of the DHFR gene, and, concomitantly, multiple copies of other DNA comprising the expression vectors, such as the DNA encoding the antibody or antibody fragment. This amplification technique can be used with any otherwise suitable host, e.g., ATCC No. CCL61 CHO-K1, notwithstanding the presence of endogenous DHFR if, for example, a mutant DHFR gene that is highly resistant to Mtx is employed (EP 117,060). Alternatively, host cells (particularly wild-type hosts that contain endogenous DHFR) transformed or co-transformed with DNA sequences encoding the antibody or antibody fragment, wild-type DHFR protein, and another selectable marker such as aminoglycoside 3' phosphotransferase (APH) can be selected by cell growth in medium containing a selection agent for the selectable marker such as an aminoglycosidic antibiotic, e.g., kanamycin, neomycin, or G418. See U.S. Pat. No. 4,965,199.

A suitable selection gene for use in yeast is the *trp*1 gene present in the yeast plasmid YRp7. Stinchcomb *et al.*, Nature, 282: 39 (1979); Kingsman *et al.*, Gene, 7: 141 (1979); or Tschemper *et al.*, Gene, 10: 157 (1980). The *trp*1 gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1. Jones, Genetics, 85: 12 (1977). The presence of the <u>trp</u>1 lesion in the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of tryptophan. Similarly, *Leu*2-deficient yeast strains (ATCC 20,622 or 38,626) are complemented by known plasmids bearing the *Leu*2 gene.

#### (iv) Promoter Component

Expression vectors usually contain a promoter that is recognized by the host organism and is operably linked to the antibody or antibody fragment nucleic acid. Promoters are untranslated sequences located upstream (5') to the start codon of a structural gene (generally within about 100 to 1000 bp) that control the transcription and translation of a particular nucleic acid sequence, such as the antibody or antibody fragment encoding sequence, to which they are operably linked. Such promoters typically fall into two classes, inducible and constitutive. Inducible promoters are promoters that initiate increased levels of transcription from DNA under their control in response to some change in culture conditions, e.g. the presence or absence of a nutrient or a change in temperature. At this time a large number of promoters recognized by a variety of potential host cells are well known.

Promoters suitable for use with prokaryotic hosts include the β-lactamase and lactose promoter

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systems (Chang et al., Nature, 275: 615 (1978); and Goeddel et al., Nature, 281: 544 (1979)), alkaline phosphatase, a tryptophan (trp) promoter system (Goeddel, Nucleic Acids Res., 8: 4057 (1980) and EP 36,776) and hybrid promoters such as the tac promoter (deBoer et al., Proc. Natl. Acad. Sci. USA, 80: 21-25 (1983)). However, other known bacterial promoters are suitable. Their nucleotide sequences have been published, thereby enabling a skilled worker to operably ligate them to DNA encoding the antibody or antibody fragment (Siebenlist et al., Cell, 20: 269 (1980)) using linkers or adaptors to supply any required restriction sites. Promoters for use in bacterial systems also generally will contain a Shine-Dalgarno (S.D.) sequence operably linked to the DNA encoding the antibody or antibody fragment.

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Suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglycerate kinase (Hitzeman et al., J. Biol. Chem., 255: 2073 (1980)) or other glycolytic enzymes (Hess et al., J. Adv. Enzyme Reg., 7: 149 (1968); and Holland, Biochemistry, 17: 4900 (1978)), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are the promoter regions for alcohol dehydrogenase 2, isocytochrome C, acid phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate dehydrogenase, and enzymes responsible for maltose and galactose utilization. Suitable vectors and promoters for use in yeast expression are further described in Hitzeman et al., EP 73,657A. Yeast enhancers also are advantageously used with yeast promoters.

Promoter sequences are known for eukaryotes. Virtually all eukaryotic genes have an AT-rich region located approximately 25 to 30 bases upstream from the site where transcription is initiated. Another sequence found 70 to 80 bases upstream from the start of transcription of many genes is a CXCAAT region where X may be any nucleotide. At the 3' end of most eukaryotic genes is an AATAAA sequence that may be the signal for addition of the poly A tail to the 3' end of the coding sequence. All of these sequences are suitably inserted into mammalian expression vectors.

Vector driven transcription of antibody or antibody fragment encoding DNA in mammalian host cells can be controlled by promoters obtained from the genomes of viruses such as polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and most preferably Simian Virus 40 (SV40), from heterologous mammalian promoters, e.g. the actin promoter or an immunoglobulin promoter, and from heat-shock promoters, provided such promoters are compatible with the host cell systems.

The early and late promoters of the SV40 virus are conveniently obtained as an SV40 restriction fragment that also contains the SV40 viral origin of replication. Fiers et al., Nature, 273: 113 (1978); Mulligan and Berg, Science, 209: 1422-1427 (1980); Pavlakis et al., Proc. Natl. Acad. Sci. USA, 78: 7398-7402 (1981). The immediate early promoter of the human cytomegalovirus is conveniently obtained as a HindIII E restriction fragment. Greenaway et al., Gene, 18: 355-360 (1982). A system for expressing DNA

in mammalian hosts using the bovine papilloma virus as a vector is disclosed in U.S. 4,419,446. A modification of this system is described in U.S. 4,601,978. See also Gray et al., Nature, 295: 503-508 (1982) on expressing cDNA encoding immune interferon in monkey cells, Reyes et al., Nature, 297: 598-601 (1982) on expression of human -interferon cDNA in mouse cells under the control of a thymidine kinase promoter from herpes simplex virus, Canaani and Berg, Proc. Natl. Acad. Sci. USA, 79: 5166-5170 (1982) on expression of the human interferon 1 gene in cultured mouse and rabbit cells, and Gorman et al., Proc. Natl. Acad. Sci. USA, 79: 6777-6781 (1982) on expression of bacterial CAT sequences in CV-1 monkey kidney cells, chicken embryo fibroblasts, Chinese hamster ovary cells, HeLa cells, and mouse NIH-3T3 cells using the Rous sarcoma virus long terminal repeat as a promoter.

### (v) Enhancer Element Component

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Transcription of a DNA encoding antibody or antibody fragment by higher eukaryotic host cells is often increased by inserting an enhancer sequence into the vector. Enhancers are cisacting elements of DNA, usually about from 10-300 bp, that act on a promoter to increase its transcription. Enhancers are relatively orientation and position independent having been found 5' (Laimins et al., Proc. Natl. Acad. Sci. USA, 78: 993 (1981)) and 3' (Lusky et al., Mol. Cell Bio., 3: 1108 (1983)) to the transcription unit, within an intron (Banerji et al., Cell, 33: 729 (1983)) as well as within the coding sequence itself (Osborne et al., Mol. Cell Bio., 4: 1293 (1984)). Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, -fetoprotein and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. See also Yaniv, Nature, 297: 17-18 (1982) on enhancing elements for activation of eukaryotic promoters. The enhancer may be spliced into the vector at a position 5' or 3' to the antibody or antibody fragment DNA, but is preferably located at a site 5' from the promoter.

## (vi) Transcription Termination Component

Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human, or nucleated cells from other multicellular organisms) can also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3' untranslated regions of eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding the antibody or antibody fragment. The 3' untranslated regions also include transcription termination sites.

Suitable vectors containing one or more of the above listed components and the desired coding and control sequences are constructed by standard ligation techniques. Isolated plasmids or DNA fragments are cleaved, tailored, and religated in the form desired to generate the plasmids required.

For analysis to confirm correct sequences in plasmids constructed, the ligation mixtures are used to transform E. coli K12 strain 294 (ATCC 31,446) and successful transformants selected by ampicillin or

tetracycline resistance where appropriate. Plasmids from the transformants are prepared, analyzed by restriction endonuclease digestion, and/or sequenced by the method of Messing et al., Nucleic Acids Res., 9: 309 (1981) or by the method of Maxam et al., Methods in Enzymology, 65: 499 (1980).

Particularly useful in the practice of this invention are expression vectors that provide for the transient expression in mammalian cells of DNA encoding the antibody or antibody fragment. In general, transient expression involves the use of an expression vector that is able to replicate efficiently in a host cell, such that the host cell accumulates many copies of the expression vector and, in turn, synthesizes high levels of a desired polypeptide encoded by the expression vector.

Other methods, vectors, and host cells suitable for adaptation to the synthesis of the antibody or antibody fragment in recombinant vertebrate cell culture are described in Gething et al., Nature, 293: 620-625 (1981); Mantei et al., Nature, 281: 40-46 (1979); Levinson et al., EP 117,060; and EP 117,058. A particularly useful plasmid for mammalian cell culture expression of the IgE peptide antagonist is pRK5 (EP pub. no. 307,247) or pSV16B (PCT pub. no. WO 91/08291 published 13 June 1991).

## C. Selection and Transformation of Host Cells

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Suitable host cells for cloning or expressing the vectors herein are the prokaryote, yeast, or higher eukaryote cells described above. Suitable prokaryotes include eubacteria, such as Gram-negative or Gram-positive organisms, for example, E. coli, Bacilli such as B. subtilis, Pseudomonas species such as P. aeruginosa, Salmonella typhimurium, or Serratia marcescens. One preferred E. coli cloning host is E. coli 294 (ATCC 31,446), although other strains such as E. coli B, E. coli 1776 (ATCC 31,537), and E. coli W3110 (ATCC 27,325) are suitable. These examples are illustrative rather than limiting. Preferably the host cell should secrete minimal amounts of proteolytic enzymes. In a preferred embodiment, the E. coli strain 49D6 is used as the expression host as described in the Examples below. Review articles describing the recombinant production of antibodies in bacterial host cells include Skerra et al., Curr. Opinion in Immunol., 5: 256 (1993) and Pluckthun, Immunol. Revs., 130: 151 (1992).

In addition to prokaryotes, eukaryotic microbes such as filamentous fungi or yeast are suitable hosts for vectors containing antibody or antibody fragment DNA. Saccharomyces cerevisiae, or common baker's yeast, is the most commonly used among lower eukaryotic host microorganisms. However, a number of other genera, species, and strains are commonly available and useful herein, such as S. pombe (Beach and Nurse, Nature, 290: 140 (1981)), Kluyveromyces lactis (Louvencourt et al., J. Bacteriol., 737 (1983)), yarrowia (EP 402,226), Pichia pastoris (EP 183,070), Trichoderma reesia (EP 244,234), Neurospora crassa (Case et al., Proc. Natl. Acad. Sci. USA, 76: 5259-5263 (1979)), and Aspergillus hosts such as A. nidulans (Ballance et al., Biochem. Biophys. Res. Commun., 112: 284-289 (1983); Tilburn et al., Gene, 26: 205-221 (1983); Yelton et al., Proc. Natl. Acad. Sci. USA, 81: 1470-1474 (1984)) and A. niger (Kelly and Hynes, EMBO J., 4: 475-479 (1985)).

Host cells derived from multicellular organisms can also be used in the recombinant production of antibody or antibody fragment. Such host cells are capable of complex processing and glycosylation activities. In principle, any higher eukaryotic cell culture is workable, whether from vertebrate or

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invertebrate culture. Examples of invertebrate cells include plant and insect cells. Numerous baculoviral strains and variants and corresponding permissive insect host cells from hosts such as Spodoptera frugiperda (caterpillar), Aedes aegypti (mosquito), Aedes albopictus (mosquito), Drosophila melanogaster (fruitfly), and Bombyx mori host cells have been identified. See, e.g., Luckow et al., Bio/Technology, 6: 47-55 (1988); Miller et al., in Genetic Engineering, Setlow, J.K. et al., 8: 277-279 (Plenum Publishing, 1986), and Maeda et al., Nature, 315: 592-594 (1985). A variety of such viral strains are publicly available, e.g., the L-1 variant of Autographa californica NPV and the Bm-5 strain of Bombyx mori NPV, and such viruses may be used as the virus herein according to the present invention, particularly for transfection of Spodoptera frugiperda cells.

Plant cell cultures of cotton, corn, potato, soybean, petunia, tomato, and tobacco can be utilized as hosts. Typically, plant cells are transfected by incubation with certain strains of the bacterium Agrobacterium tumefaciens, which has been previously manipulated to contain the antibody or antibody fragment DNA. During incubation of the plant cell culture with A. tumefaciens, the DNA encoding antibody or antibody fragment is transferred to the plant cell host such that it is transfected, and will, under appropriate conditions, express the antibody or antibody fragment DNA. In addition, regulatory and signal sequences compatible with plant cells are available, such as the nopaline synthase promoter and polyadenylation signal sequences. Depicker et al., J. Mol. Appl. Gen., 1: 561 (1982). In addition, DNA segments isolated from the upstream region of the T-DNA 780 gene are capable of activating or increasing transcription levels of plant-expressible genes in recombinant DNA-containing plant tissue. See EP 321,196 published 21 June 1989.

Vertebrate cell culture is preferred for the recombinant production of full length antibodies. The propagation of vertebrate cells in culture (tissue culture) has become a routine procedure in recent years (Tissue Culture, Academic Press, Kruse and Patterson, editors (1973)). Examples of useful mammalian host cell lines are monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham et al., J. Gen Virol., 36: 59 (1977)); baby hamster kidney cells (BHK, ATCC CCL 10); Chinese hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, 77: 4216 (1980)); mouse sertoli cells (TM4, Mather, Biol. Reprod., 23: 243-251 (1980)); monkey kidney cells (CV1 ATCC CCL 70); African green monkey kidney cells (VERO-76, ATCC CRL-1587); human cervical carcinoma cells (HELA, ATCC CCL 2); canine kidney cells (MDCK, ATCC CCL 34); buffalo rat liver cells (BRL 3A, ATCC CRL 1442); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); mouse mammary tumor (MMT 060562, ATCC CCL51); TRI cells (Mather et al., Annals N.Y. Acad. Sci., 383: 44-68 (1982)); MRC 5 cells; FS4 cells; and a human hepatoma cell line (Hep G2). Preferred host cells are human embryonic kidney 293 and Chinese hamster ovary cells. Myeloma cells that do not otherwise produce immunoglobulin protein are also useful host cells for the recombinant production of full length antibodies.

Host cells are transfected and preferably transformed with the above-described expression or cloning vectors of this invention and cultured in conventional nutrient media modified as appropriate for

inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences.

Transfection refers to the taking up of an expression vector by a host cell whether or not any coding sequences are in fact expressed. Numerous methods of transfection are known to the ordinarily skilled artisan, for example, CaPO<sub>4</sub> precipitation and electroporation. Successful transfection is generally recognized when any indication of the operation of this vector occurs within the host cell.

Transformation means introducing DNA into an organism so that the DNA is replicable, either as an extrachromosomal element or by chromosomal integrant. Depending on the host cell used, transformation is done using standard techniques appropriate to such cells. The calcium treatment employing calcium chloride, as described in section 1.82 of Sambrook et al., supra, is generally used for prokaryotes or other cells that contain substantial cell-wall barriers. Infection with Agrobacterium tumefaciens is used for transformation of certain plant cells, as described by Shaw et al., Gene, 23: 315 (1983) and WO 89/05859 published 29 June 1989. For mammalian cells without such cell walls, the calcium phosphate precipitation method described in sections 16.30-16.37 of Sambrook et al., supra, is preferred. General aspects of mammalian cell host system transformations have been described by Axel in U.S. 4,399,216 issued 16 August 1983. Transformations into yeast are typically carried out according to the method of Van Solingen et al., J. Bact., 130: 946 (1977) and Hsiao et al., Proc. Natl. Acad. Sci. (USA), 76: 3829 (1979). However, other methods for introducing DNA into cells such as by nuclear injection, electroporation, or by protoplast fusion may also be used.

#### D. Culturing the Host Cells

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Prokaryotic cells used to produce the antibody or antibody fragment are cultured in suitable media as described generally in Sambrook et al., supra.

The mammalian host cells used to produce the antibody or antibody fragment can be cultured in a variety of media. Commercially available media such as Ham's F10 (Sigma), Minimal Essential Medium ((MEM), Sigma), RPMI-1640 (Sigma), and Dulbecco's Modified Eagle's Medium ((DMEM), Sigma) are suitable for culturing the host cells. In addition, any of the media described in Ham and Wallace, Meth. Enz., 58: 44 (1979), Barnes and Sato, Anal. Biochem., 102: 255 (1980), U.S. 4,767,704; 4,657,866; 4,927,762; or 4,560,655; WO 90/03430; WO 87/00195; U.S. Pat. Re. 30,985; or U.S. 5,122,469, the disclosures of all of which are incorporated herein by reference, may be used as culture media for the host cells. Any of these media may be supplemented as necessary with hormones and/or other growth factors (such as insulin, transferrin, or epidermal growth factor), salts (such as sodium chloride, calcium, magnesium, and phosphate), buffers (such as HEPES), nucleosides (such as adenosine and thymidine), antibiotics (such as Gentamycin TM drug), trace elements (defined as inorganic compounds usually present at final concentrations in the micromolar range), and glucose or an equivalent energy source. Any other necessary supplements may also be included at appropriate concentrations that would be known to those skilled in the art. The culture conditions, such as temperature, pH, and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

The host cells referred to in this disclosure encompass cells in in vitro culture as well as cells that

are within a host animal.

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### E. <u>Detecting Gene Amplification/Expression</u>

Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, northern blotting to quantitate the transcription of mRNA (Thomas, Proc. Natl. Acad. Sci. USA, 77: 5201-5205 (1980)), dot blotting (DNA analysis), or in situ hybridization, using an appropriately labeled probe, based on the sequences provided herein. Various labels may be employed, most commonly radioisotopes, particularly <sup>32</sup>P. However, other techniques may also be employed, such as using biotin-modified nucleotides for introduction into a polynucleotide. The biotin then serves as the site for binding to avidin or antibodies, which may be labeled with a wide variety of labels, such as radionuclides, fluorescers, enzymes, or the like. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes. The antibodies in turn may be labeled and the assay may be carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected.

Gene expression, alternatively, may be measured by immunological methods, such as immunohistochemical staining of tissue sections and assay of cell culture or body fluids, to quantitate directly the expression of gene product. With immunohistochemical staining techniques, a cell sample is prepared, typically by dehydration and fixation, followed by reaction with labeled antibodies specific for the gene product, where the labels are usually visually detectable, such as enzymatic labels, fluorescent labels, luminescent labels, and the like. A particularly sensitive staining technique suitable for use in the present invention is described by Hsu et al., Am. J. Clin. Path., 75: 734-738 (1980).

#### F. Purification of the Antibody or Antibody Fragment

In the case of a host cell secretion system, the antibody or antibody fragment is recovered from the culture medium. Alternatively, the antibody can be produced intracellularly, or produced in the periplasmic space of a bacterial host cell. If the antibody is produced intracellularly, as a first step, the host cells are lysed, and the resulting particulate debris is removed, for example, by centrifugation or ultrafiltration. Carter et al., Bio/Technology 10:163-167 (1992) describe a procedure for isolating antibodies which are secreted to the periplasmic space of E. coli. Briefly, cell paste is thawed in the presence of sodium acetate (pH 3.5), EDTA, and phenylmethylsulfonylfluoride (PMSF) over about 30 min. Cell debris can be removed by centrifugation. Where the antibody is secreted into the medium, supernatants from such expression systems are generally first concentrated using a commercially available protein concentration filter, for example, an Amicon or Millipore Pellicon ultrafiltration unit. A protease inhibitor such as PMSF may be included in any of the foregoing steps to inhibit proteolysis and antibiotics may be included to prevent the growth of adventitious contaminants.

The antibody composition prepared from the cells can be purified using, for example, hydroxylapatite chromatography, gel electrophoresis, dialysis, and affinity chromatography, with affinity chromatography being the preferred purification technique. The suitability of protein A as an affinity ligand

depends on the species and isotype of any immunoglobulin Fc domain that is present in the antibody. Protein A can be used to purify antibodies that are based on human γ1, γ2, or γ4 heavy chains (Lindmark et al., J. Immunol. Meth. 62:1-13 (1983)). Protein G is recommended for all mouse isotypes and for human γ3 (Guss et al., EMBO J. 5:15671575 (1986)). The matrix to which the affinity ligand is attached is most often agarose, but other matrices are available. Mechanically stable matrices such as controlled pore glass or poly(styrenedivinyl)benzene allow for faster flow rates and shorter processing times than can be achieved with agarose. Where the antibody comprises a CH3 domain, the Bakerbond ABX<sup>TM</sup>resin (J. T. Baker, Phillipsburg, NJ) is useful for purification. Other techniques for protein purification such as fractionation on an ion-exchange column, ethanol precipitation, Reverse Phase HPLC, chromatography on silica, chromatography on heparin Sepharose<sup>TM</sup> chromatography on an anion or cation exchange resin (such as a polyaspartic acid column), chromatofocusing, SDS-PAGE, and ammonium sulfate precipitation are also available depending on the antibody to be recovered.

Following any preliminary purification step(s), the mixture comprising the antibody of interest and contaminants may be subjected to low pH hydrophobic interaction chromatography using an elution buffer at a pH between about 2.5-4.5, preferably performed at low salt concentrations (e.g. from about 0-0.25M salt).

## G. Production of Antibody Fragments

Various techniques have been developed for the production of the humanized antibody fragments of the invention, including Fab, Fab', Fab'-SH, or F(ab') 2 fragments. Traditionally, these fragments were derived via proteolytic digestion of intact antibodies (see, e.g., Morimoto et al., Journal of Biochemical and Biophysical Methods 24:107-117 (1992) and Brennan et al., Science, 229:81 (1985)). However, these fragments can now be produced directly by recombinant host cells. For example, Fab'-SH fragments can be directly recovered from E. coli and chemically coupled to form F(ab') 2 fragments (Carter et al., Bio/Technology, 10:163-167 (1992)). According to another approach, F(ab') 2 fragments can be isolated directly from recombinant host cell culture. Other techniques for the production of antibody fragments will be apparent to the skilled practitioner.

## 5. Uses of Anti-IL-8 Antibodies

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#### A. Diagnostic Uses

For diagnostic applications requiring the detection or quantitation of IL-8, the antibodies or antibody fragments of the invention typically will be labeled with a detectable moiety. The detectable moiety can be any one which is capable of producing, either directly or indirectly, a detectable signal. For example, the detectable moiety can be a radioisotope, such as <sup>3</sup>H, <sup>14</sup>C, <sup>32</sup>P, <sup>35</sup>S, or <sup>125</sup>I; a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin; radioactive isotopic labels, such as, e.g., <sup>125</sup>I, <sup>32</sup>P, <sup>14</sup>C, or <sup>3</sup>H; or an enzyme, such as alkaline phosphatase, betagalactosidase, or horseradish peroxidase.

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Any method known in the art for separately conjugating the antibody or antibody fragment to the detectable moiety can be employed, including those methods described by Hunter et al., Nature 144:945 (1962); David et al., Biochemistry 13:1014 (1974); Pain et al., J. Immunol. Meth. 40:219 (1981); and Nygren, J. Histochem. and Cytochem. 30:407 (1982).

The antibodies and antibody fragments of the present invention can be employed in any known assay method, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays. For example, see Zola, Monoclonal Antibodies: A Manual of Techniques, pp. 147-158 (CRC Press, Inc., 1987).

Competitive binding assays rely on the ability of a labeled standard (which can be a IL-8 or an immunologically reactive portion thereof) to compete with the test sample analyte (IL-8) for binding with a limited amount of antibody or antibody fragment. The amount of IL-8 in the test sample is inversely proportional to the amount of standard that becomes bound to the antibodies. To facilitate determining the amount of standard that becomes bound, the antibodies or antibody fragments generally are insolubilized before or after the competition, so that the standard and analyte that are bound to the antibodies can conveniently be separated from the standard and analyte which remain unbound.

Sandwich assays involve the use of two antibodies, each capable of binding to a different antigenic portion, or epitope, of the protein (IL-8) to be detected. In a sandwich assay, the test sample analyte is bound by a first antibody which is immobilized on a solid support, and thereafter a second antibody binds to the analyte, thus forming an insoluble three-part complex (U.S. Patent No. 4,376,110). The second antibody can itself be labeled with a detectable moiety (direct sandwich assays) or can be measured using an anti-immunoglobulin antibody that is labeled with a detectable moiety (indirect sandwich assay). For example, one type of sandwich assay is an ELISA assay, in which case the detectable moiety is an enzyme (e.g., horseradish peroxidase).

IL-8 antibodies and antibody fragments also are useful for the affinity purification of IL-8 from recombinant cell culture or natural sources. For example, these antibodies can be fixed to a solid support by techniques well known in the art so as to purify IL-8 from a source such as culture supernatant or tissue.

## B. Therapeutic Compositions and Administration of Anti-IL-8 Antibody

The humanized anti-IL-8 antibodies and antibody fragments of the invention are useful in the treatment of inflammatory disorders, such as adult respiratory distress syndrome (ARDS), hypovolemic shock, ulcerative colitis, and rheumatoid arthritis.

Therapeutic formulations of the humanized anti-IL-8 antibodies and antibody fragments are prepared for storage by mixing the antibody or antibody fragment having the desired degree of purity with optional physiologically acceptable carriers, excipients, or stabilizers (Remington's Pharmaceutical Sciences, supra), in the form of lyophilized cake or aqueous solutions. Acceptable carriers, excipients or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid; low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin, or immunoglobulins;

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hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or nonionic surfactants such as Tween, Pluronics or polyethylene glycol (PEG).

The humanized anti-IL-8 mAb or antibody fragment to be used for in vivo administration must be sterile. This is readily accomplished by filtration through sterile filtration membranes, prior to or following lyophilization and reconstitution. The humanized anti-IL-8 mAb or antibody fragment ordinarily will be stored in lyophilized form or in solution.

Therapeutic humanized anti-IL-8 mAb or antibody fragment compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

The route of humanized anti-IL-8 mAb or antibody fragment administration is in accord with known methods, e.g., inhalation, injection or infusion by intravenous, intraperitoneal, intracerebral, intramuscular, intraocular, intraarterial, or intralesional routes, by enema or suppository, or by sustained release systems as noted below. Preferably the antibody is given systemically or at a site of inflammation.

Suitable examples of sustained-release preparations include semipermeable polymer matrices in the form of shaped articles, e.g. films, or microcapsules. Sustained release matrices include polyesters, hydrogels, polylactides (U.S. 3,773,919, EP 58,481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman et al., Biopolymers 22:547 (1983)), poly (2-hydroxyethyl-methacrylate) (Langer et al., J. Biomed. Mater. Res. 15:167 (1981) and Langer, Chem. Tech. 12:98 (1982)), ethylene vinyl acetate (Langer et al., supra) or poly-D-(-)-3-hydroxybutyric acid (EP 133,988). Sustained-release humanized anti-IL-8 antibody or antibody fragment compositions also include liposomally entrapped antibody or antibody fragment. Liposomes containing an antibody or antibody fragment are prepared by methods known per se: DE 3,218,121; Epstein et al., Proc. Natl. Acad. Sci. U.S.A. 82:3688 (1985); Hwang et al., Proc. Natl. Acad. Sci. U.S.A. 77:4030 (1980); EP 52,322; EP 36,676; EP 88,046; EP 143,949; EP 142,641; Japanese patent application 83-118008; U.S. Patent Nos. 4,485,045 and 4,544,545; and EP 102,324. Ordinarily the liposomes are of the small (about 200-800 Angstroms) unilamelar type in which the lipid content is greater than about 30 mole percent cholesterol, the selected proportion being adjusted for the most efficacious antibody or antibody fragment therapy.

An "effective amount" of the humanized anti-IL-8 antibody or antibody fragment to be employed therapeutically will depend, for example, upon the therapeutic objectives, the route of administration, and the condition of the patient. Accordingly, it will be necessary for the therapist to titer the dosage and modify the route of administration as required to obtain the optimal therapeutic effect. Typically, the clinician will administer the humanized anti-IL-8 antibody or antibody fragment until a dosage is reached that achieves the desired effect. The progress of this therapy is easily monitored by conventional assays.

In the treatment and prevention of an inflammatory disorder with a humanized anti-IL-8 antibody

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or antibody fragment of the invention, the antibody composition will be formulated, dosed, and administered in a fashion consistent with good medical practice. Factors for consideration in this context include the particular disorder being treated, the particular mammal being treated, the clinical condition of the individual patient, the cause of the disorder, the site of delivery of the antibody, the particular type of antibody, the method of administration, the scheduling of administration, and other factors known to medical practitioners. The "therapeutically effective amount" of antibody to be administered will be governed by such considerations, and is the minimum amount necessary to prevent, ameliorate, or treat the inflammatory disorder, including treating acute or chronic respiratory diseases and reducing inflammatory responses. Such amount is preferably below the amount that is toxic to the host or renders the host significantly more susceptible to infections.

As a general proposition, the initial pharmaceutically effective amount of the antibody or antibody fragment administered parenterally per dose will be in the range of about 0.1 to 50 mg/kg of patient body weight per day, with the typical initial range of antibody used being 0.3 to 20 mg/kg/day, more preferably 0.3 to 15 mg/kg/day.

As noted above, however, these suggested amounts of antibody or antibody fragment are subject to a great deal of therapeutic discretion. The key factor in selecting an appropriate dose and scheduling is the result obtained, as indicated above.

The antibody or antibody fragment need not be, but is optionally formulated with one or more agents currently used to prevent or treat the inflammatory disorder in question. For example, in rheumatoid arthritis, the antibody can be given in conjunction with a glucocorticosteroid. The effective amount of such other agents depends on the amount of antibody or antibody fragment present in the formulation, the type of disorder or treatment, and other factors discussed above. These are generally used in the same dosages and with administration routes as used hereinbefore or about from 1 to 99% of the heretofore employed dosages.

The following examples are offered by way of illustration and not by way of limitation. The disclosures of all references cited in the specification, and the disclosures of all citations in such references, are expressly incorporated herein by reference.

#### **EXAMPLES**

## A. GENERATION AND CHARACTERIZATION OF MONOCLONAL ANTIBODIES AGAINST HUMAN IL-8

Balb/c mice were immunized in each hind footpad or intraperitoneally with 10 μg of recombinant human IL-8 (produced as a fusion of (ser-IL-8)<sub>72</sub> with ubiquitin (Hebert *et al.* J. Immunology 145:3033-3040 (1990)); IL-8 is available commercially from PeproTech, Inc., Rocky Hill, NJ) resuspended in MPL/TDM (Ribi Immunochem. Research Inc., Hamilton, MT) and boosted twice with the same amount of IL-8. In these experiments, "IL-8" is intended to mean (ser-IL-8)<sub>72</sub> unless otherwise specified. A final boost of 10 μg of IL-8 was given 3 days before the fusion. Spleen cells or popliteal lymph node cells were fused with mouse myeloma P3X63Ag8U.1 (ATCC CRL1597), a non-secreting clone of the myeloma P3X63Ag8, using 35% polyethylene glycol as described before. Ten days after the fusion, culture supernatant was

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screened for the presence of monoclonal antibodies to IL-8 by ELISA.

The ELISA was performed as foll ws. Nunc 96-well immunoplates (Flow Lab, McLean, VA) were coated with 50 μl/well of 2 μg/ml IL-8 in phosphate-buffered saline (PBS) overnight at 4°C. The remaining steps were carried out at room temperature. Nonspecific binding sites were blocked with 0.5% bovine serum albumin (BSA) for 1 hour (hr). Plates were then incubated with 50 μl/well of hybridoma culture supernatants from 672 growing parental fusion wells for 1 hr, followed by the incubation with 50 μl/well of 1:1000 dilution of a 1 mg/ml stock solution of alkaline phosphatase-conjugated goat anti-mouse Ig (Tago Co., Foster City, CA) for 1 hr. The level of enzyme-linked antibody bound to the plate was determined by the addition of 100 μl/well of 0.5 mg/ml of r-nitrophenyl phosphate in sodium bicarbonate buffer, pH 9.6. The color reaction was measured at 405 nm with an ELISA plate reader (Titertrek Multiscan, Flow Lab, McLean, VA). Between each step, plates were washed three times in PBS containing 0.05% Tween 20.

Culture supernatants which promoted 4-fold more binding of IL-8 than did control medium were selected as positives. According to this criterion, 16 of 672 growing parental fusion wells (2%) were positive. These positive hybridoma cell lines were cloned at least twice by using the limiting dilution technique.

Seven of the positive hybridomas were further characterized as follows. The isotypes of the monoclonal antibodies were determined by coating Nunc 96-well immunoplates (Flow Lab, McLean, VA) with IL-8 overnight, blocking with BSA, incubating with culture supernatants followed by the addition of predetermined amount of isotype-specific alkaline phosphatase-conjugated goat anti-mouse Ig (Fisher Biotech, Pittsburgh, PA). The level of conjugated antibodies bound to the plate was determined by the addition of r-nitrophenyl phosphate as described above.

All the monoclonal antibodies tested belonged to either IgG<sub>1</sub> or IgG<sub>2</sub> immunoglobulin isotype. Ascites fluid containing these monoclonal antibodies had antibody titers in the range of 10,000 to 100,000 as determined by the reciprocal of the dilution factor which gave 50% of the maximum binding in the ELISA.

To assess whether these monoclonal antibodies bound to the same epitopes, a competitive binding ELISA was performed. At a ratio of biotinylated mAb to unlabeled mAb of 1:100, the binding of biotinylated mAb 5.12.14 was significantly inhibited by its homologous mAb but not by mAb 4.1.3, while the binding of biotinylated mAb 4.1.3 was inhibited by mAb 4.1.3 but not by mAb 5.12.14. Monoclonal antibody 5.2.3 behaved similarly to mAb 4.1.3, while monoclonal antibodies 4.8 and 12.3.9 were similar to mAb 5.12.14. Thus, mAb 4.1.3 and mAb 5.2.3 bind to a different epitope(s) than the epitope recognized by monoclonal antibodies 12.3.9, 4.8 and 5.12.14.

Immunodot blot analysis was performed to assess antibody reactivity to IL-8 immobilized on nitrocellulose paper. All seven antibodies recognized IL-8 immobilized on paper, whereas a control mouse IgG antibody did not.

The ability of these monoclonal antibodies to capture soluble 1251-1L-8 was assessed by a

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radioimmune precipitation test (RIP). Briefly, tracer <sup>125</sup>I-IL-8 (4 x 10<sup>4</sup> cpm) was incubated with various dilutions of the monoclonal anti-IL-8 antibodies in 0.2 ml of PBS containing 0.5% BSA and 0.05% Tween 20 (assay buffer) for 1 hr at room temperature. One hundred microliters of a predetermined concentration of goat anti-mouse Ig antisera (Pel-Freez, Rogers, AR) were added and the mixture was incubated at room temperature for 1 hr. Immune complexes were precipitated by the addition of 0.5 ml of 6% polyethylene glycol (M.W. 8000) kept at 4°C. After centrifugation at 2,000 x g for 20 min at 4°C, the supernatant was removed by aspiration and the radioactivity remaining in the pellet was counted in a gamma counter. Percent specific binding was calculated as (precipitated cpm - background cpm)/ (total cpm - background cpm). Monoclonal antibodies 4.1.3, 5.2.3, 4.8, 5.12.14 and 12.3.9 captured <sup>125</sup>I-IL-8 very efficiently, while antibodies 9.2.4 and 8.9.1 were not able to capture soluble <sup>125</sup>I-IL-8 in the RIP even though they could bind to IL-8 coated onto ELISA plates (Table I).

The dissociation constants of these monoclonal antibodies were determined using a competitive binding RIP assay. Briefly, competitive inhibition of the binding each antibody to  $^{125}$ I-IL-8 (20,000-40,000 cpm per assay) by various amounts of unlabeled IL-8 was determined by the RIP described above. The dissociation constant (affinity)of each mAb was determined by using Scatchard plot analysis (Munson, et al., Anal. Biochem. 107:220 (1980)) as provided in the VersaTerm-PRO computer program (Synergy Software, Reading, PA). The  $K_d$ 's of these monoclonal antibodies (with the exception of 9.2.4. and 8.9.1) were in the range from 2 x  $10^{-8}$  to 3 x  $10^{-10}$  M. Monoclonal antibody 5.12.14 with a  $K_d$  of 3 x  $10^{-10}$  M showed the highest affinity among all the monoclonal antibodies tested (Table 3).

Table 3. Characterization of Anti-IL-8 Monoclonal Antibodies

Antibody	%Specific Binding to IL-8	K <sub>d</sub> (M)	Isotype	pl
4.1.3	58	2 X 10 <sup>-9</sup>	lgG <sub>l</sub>	4.3-6.1
5.2.3	34	2 X 10 <sup>-8</sup>	IgG <sub>1</sub>	5.2-5.6
9.2.4	1	-	IgG <sub>1</sub>	7.0-7.5
8.9.1	2	-	IgG <sub>1</sub>	6.8-7.6

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Antibody	%Specific Binding to IL-8	K <sub>d</sub> (M)	Isotype	pI
4.8	62	3 X 10 <sup>-8</sup>	IgG <sub>2a</sub>	6.1-7.1
5.12.14	98	3 X 10 <sup>-10</sup>	IgG <sub>2a</sub>	6.2-7.4
12.3.9	86	2 X 10 <sup>-9</sup>	IgG <sub>2a</sub>	6.5-7.1

To assess the ability of these monoclonal antibodies to neutralize IL-8 activity, the amount of <sup>125</sup>I-IL-8 bound to human neutrophils in the presence of various amounts of culture supernatants and purified monoclonal antibodies was measured. Neutrophils were prepared by using Mono-Poly Resolving Medium (M-PRM) (Flow Lab. Inc., McLean, VA). Briefly fresh, heparinized human blood was loaded onto M-PRM at a ratio of blood to medium, 3.5:3.0, and centrifuged at 300 x g for 30 min at room temperature. Neutrophils enriched at the middle layer were collected and washed once in PBS. Such a preparation routinely contained greater than 95% neutrophils according to the Wright's Giernsa staining. The receptor binding assay was done as follows. 50 μl of <sup>125</sup>I-IL-8 (5 ng/ml) was incubated with 50 μl of unlabeled IL-8 (100 μg/ml) or monoclonal antibodies in PBS containing 0.1% BSA for 30 min at room temperature. The mixture was then incubated with 100 μl of neutrophils (10<sup>7</sup> cells/ml) for 15 min at 37°C. The <sup>125</sup>I-IL-8 bound was separated from the unbound material by loading mixtures onto 0.4 ml of PBS containing 20% sucrose and 0.1% BSA and by centrifugation at 300 x g for 15 min. The supernatant was rem ved by aspiration and the radioactivity associated with the pellet was counted in a gamma counter.

binding of IL-8 to human neutrophils at a 1:25 molar ratio of IL-8 to mAb. On the other hand, monoclonal antibodies 9.2.4 and 8.9.1 appeared to enhance the binding of IL-8 to its receptors on human neutrophils. Since a control mouse IgG also enhanced the binding of IL-8 on neutrophils, the enhancement of IL-8 binding to its receptors by mAb 9.2.4 and 8.9.1 appears to be nonspecific. Thus, monoclonal antibodies, 4.1.3, 5.1.3, 4.8, 5.12.14, and 12.3.9 are potential neutralizing monoclonal antibodies while monoclonal

Monoclonal antibodies 4.1.3, 5.2.3, 4.8, 5.12.14, and 12.3.9 inhibited greater than 85% of the

antibodies 8.9.1 and 9.2.4 are non-neutralizing monoclonal antibodies.

The ability of the anti-IL-8 antibodies to block neutrophil chemotaxis induced by IL-8 was tested as follows. Neutrophil chemotaxis induced by IL-8 was determined using a Boyden chamber method

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(Larsen, et al. Science 243:1464 (1989)). One hundred  $\mu$ l of human neutrophils (10<sup>6</sup> cells/ml) resuspended in RPMI containing 0.1% BSA were placed in the upper chamber and 29  $\mu$ l of the IL-8 (20 nM) with or without monoclonal antibodies were placed in the lower chamber. Cells were incubated for 1 hr at 37°C. Neutrophils migrated into the lower chamber were stained with Wright's Giemsa stain and counted under the microscope (100x magnification). Approximately 10 different fields per experimental group were examined. Neutralizing monoclonal antibodies 5.12.14 and 4.1.3 blocked almost 70% of the neutrophil chemotactic activity of IL-8 at 1:10 ratio of IL-8 to mAb.

The isoelectric focusing (IEF) pattern of each mAb was determined by applying purified antibodies on an IEF polyacrylamide gel (pH 3-9, Pharmacia) using the Fast gel system (Pharmacia, Piscataway, NJ). The IEF gel was pretreated with pharmalyte containing 1% Triton X100 (Sigma, St. Louis, MO) for 10 min before loading the samples. The IEF pattern was visualized by silver staining according to the instructions from the manufacturer. All of the monoclonal antibodies had different IEF patterns, confirming that they originated from different clones. The pl values for the antibodies are listed in Table 3.

All these monoclonal antibodies bound equally well to both (ala-IL-8)77 and (ser-IL-8)72 forms of IL-8. Because IL-8 has greater than 30% sequence homology with certain other members of the platelet factor 4 (PF4) family of inflammatory cytokines such as β-TG (Van Damme et al., Eur. J. Biochem. 181:337(1989); Tanaka et al., FEB 236(2):467 (1988)) and PF4 (Deuel et al., Proc. Natl. Acad. Sci. U.S.A. 74:2256 (1977)), they were tested for possible cross reactivity to β-TG and PF4, as well as to another neutrophil activating factor, C5a. No detectable binding to any of these proteins was observed, with the exception of mAb 4.1.3, which had a slight cross reactivity to β-TG.

One of the antibodies, mAb 5.12.14, was further studied to determine whether it could block the IL-8 mediated release of elastase by neutrophils. Briefly, human neutrophils were resuspended in Hanks balanced salt solution (Gibco, Grand Island, NY) containing 1.0% BSA, Fraction V (Sigma, St. Louis, MO), 2 mg/ml alpha-D-glucose (Sigma), 4.2 mM sodium bicarbonate (Sigma) and 0.01 M HEPES, pH 7.1 (JRH Bioscience, Lenexa, KS). A stock of cytochalasin B (Sigma) was prepared (5 mg/ml in dimethylsulfoxide (Sigma) and stored at 2-8°C. Cytochalasin B was added to the neutrophil preparation to produce a final concentration of 5 µg/ml, and incubated for 15 min at 37°C. Human IL-8 was incubated with mAb 5.12.14 (20 µl), or a negative control antibody, in 1 ml polypropylene tubes (DBM Scientific, San Fernando, CA) for 30 min at 37°C. The final assay concentrations of IL-8 were 50 and 500 nM. The monoclonal antibodies were diluted to produce the following ratios (IL-8:Mab): 1:50, 1:10, 1:2, 1:1, and 1:0.25. Cytochalasin B-treated neutrophils were added (100 µl/tube) and incubated for 2 hours at 25°C. The tubes were centrifuged (210 X g, 2-8°C) for 10 min, and supernatants were transferred to 96 well tissue culture plates (30 µl/well). Elastase substrate stock, 10 mM methoxysuccinyl-alanyl-alanyl-propyl-valyl-pnitroanilide (Calbiochem, La Jolla, CA) in DMSO was prepared and stored at 2-8°C. Elastase substrate solution (1.2 mM substrate, 1.2 M NaCl (Mallinckrodt, Paris, Kentucky), 0.12 M HEPES pH 7.2 in distilled water) was added (170 µl/well) to the supernatants and incubated for 0.5 to 2 hours at 37°C (until control

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O.D. of 1.0 was reached). Absorbance was measured at 405 nm (SLT 340 ATTC plate reader, SLT Lab Instruments, Austria).

The results are shown in Figure 1. At a 1:1 ratio of IL-8 to mAb 5.12.14, the antibody was able to effectively block the release of elastase from neutrophils.

The hybridoma producing antibody 5.12.14 was deposited on February 15, 1993 with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, U.S.A. (ATCC) and assigned ATTC Accession No. HB 11553.

## B. <u>GENERATION AND CHARACTERIZATION OF MONOCLONAL ANTIBODIES AGAINST</u> RABBIT IL-8

Antibodies against rabbit IL-8 were generated in essentially the same process as anti-human IL-8 antibodies using rabbit IL-8 as immunogen (kindly provided by C. Broaddus; see also Yoshimura et al. J. Immunol. 146:3483 (1991)). The antibody was characterized as described above for binding to other cytokines coated onto ELISA plates; no measurable binding was found to MGSA, fMLP, C5a, b-TG, TNF, PF4, or IL-1.

The hybridoma producing antibody 6G4.2.5 was deposited on September 28, 1994, with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, U.S.A. (ATCC) and assigned ATTC Accession No. HB 11722.

Recombinant human-murine chimeric Fabs for 5.12.14 and 6G4.2.5 were constructed as described below. A chimeric 6G.4.25 Fab is compared with a chimeric 5.12.14 Fab in detail below.

### 1. <u>INHIBITION OF IL-8 BINDING TO HUMAN NEUTROPHILS BY 5.12.14-FAB AND 6G4</u> 2.5-FAB

The ability of the two chimeric Fabs, 5.12.14-Fab and 6G4.2.5-Fab, to efficiently bind IL-8 and prevent IL-8 from binding to IL-8 receptors on human neutrophils was determined by performing a competition binding assay which allows the calculation of the IC<sub>50</sub> - concentration required to achieve 50% inhibition of IL-8 binding.

Human neutrophils (5 X 10<sup>5</sup>), were incubated for 1 hour at 4°C with 0.5nM <sup>125</sup>I-IL-8 in the presence of various concentrations (0 to 300 nM) of 5.12.14-Fab, 6G4.2.5-Fab, an isotype control (4D5-Fab) or unlabeled IL-8. After the incubation, the unbound <sup>125</sup>I-IL-8 was removed by centrifugation through a solution of 20% sucrose and 0.1% bovine serum albumin in phosphate buffered saline and the amount of <sup>125</sup>I-IL-8 bound to the cells was determined by counting the cell pellets in a gamma counter. Figure 2 demonstrates the inhibition of <sup>125</sup>I-IL-8 binding to neutrophils by unlabeled IL-8. Figure 3 demonstrates that a negative isotype matched Fab does not inhibit the binding of <sup>125</sup>I-IL-8 to human neutrophils. Both the anti-IL-8 Fabs, 5.12.14 Fab (Figure 4) and 6G.4.25 Fab (Figure 5) were able to inhibit the binding of <sup>125</sup>I-IL-8 to human neutrophils with an average IC<sub>50</sub> of 1.6 nM and 7.5 nM, respectively.

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### 2. <u>INHIBITION OF IL-8-MEDIATED NEUTROPHIL CHEMOTAXIS BY 5.12.14-FAB AND 6G4.2.5-FAB</u>

Human neutrophils were isolated, counted and resuspended at 5 x  $10^6$  cells/ml in Hank's balanced salt solution (abbreviated HBSS; without calcium and magnesium) with 0.1% bovine serum albumin. The neutrophils were labeled by adding calcein AM (Molecular Probe, Eugene, OR) at a final concentration of 2.0  $\mu$ M. Following a 30 minute incubation at 37°C, cells were washed twice with HBSS-BSA and resuspended at 5 x  $10^6$  cells/ml.

Chemotaxis experiments were carried out in a Neuro Probe (Cabin John, MD) 96-well chamber, model MBB96. Experimental samples (buffer only control, IL-8 alone or IL-8 + Fabs) were loaded in a Polyfiltronics 96-well View plate (Neuro Probe Inc.) placed in the lower chamber. 100 µl of the calcein AM-labeled neutrophils were added to the upper chambers and allowed to migrate through a 5 micrometer porosity PVP free polycarbonate framed filter (Neuro Probe Inc.) toward the bottom chamber sample. The chemotaxis apparatus was then incubated for 40 to 60 minutes at 37°C with 5% CO<sub>2</sub>. At the end of the incubation, neutrophils remaining in the upper chamber were aspirated and upper chambers were washed three times with PBS. Then the polycarbonate filter was removed, non-migrating cells were wiped off with a squeegee wetted with PBS, and the filter was air dried for 15 minutes.

The relative number of neutrophils migrating through the filter (Neutrophil migration index) was determined by measuring fluorescence intensity of the filter and the fluorescence intensity of the contents of the lower chamber and adding the two values together. Fluorescence intensity was measured with a CytoFluor 2300 fluorescent plate reader (Millipore Corp. Bedford, MA) configured to read a Corning 96-well plate using the 485-20 nm excitation filter and a 530-25 emission filter, with the sensitivity set at 3.

The results are shown in Figures 6 and 7. Figure 6 demonstrates the inhibition of human IL-8 mediated neutrophil chemotaxis by chimeric 6G4.2.5 and 5.12.14 Fabs. Figure 7 demonstrates the relative abilities of chimeric 6G4.2.5 and 5.12.14 Fabs to inhibit rabbit IL-8 mediated neutrophil chemotaxis.

### 3. <u>INHIBITION OF IL-8-MEDIATED NEUTROPHIL ELASTASE RELEASE BY VARIOUS</u> CONCENTRATIONS OF 6G4.2.5 AND 5.12.14 FABS

Blood was drawn from healthy male donors into heparinized syringes. Neutrophils were isolated by dextran sedimentation, centrifugation over Lymphocyte Separation Medium (Organon Teknika, Durham, NC), and hypotonic lysis of contaminating red blood cells as described by Berman *et al.* (J. Cell Biochem. 52:183 (1993)). The final neutrophil pellet was suspended at a concentration of 1 x 10<sup>7</sup> cells/ml in assay buffer, which consisted of Hanks Balanced Salt Solution (GIBCO, Grand Island, NY) supplemented with 1.0% BSA (fraction V, Sigma, St. Louis, MO), 2 mg/ml glucose, 4.2 mM sodium bicarbonate, and 0.01 M HEPES, pH 7.2. The neutrophils were stored at 4°C for not longer than 1 hr.

IL-8 (10 µl) was mixed with anti-IL-8 Fab, an isotype control Fab, or buffer (20 µl) in 1 ml polypropylene tubes and incubated in a 37°C water bath for 30 min. IL-8 was used at final concentrations ranging from 0.01 to 1000 nM in dose response studies (Figure 8) and at a final concentration of 100 nM in

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the experiments addressing the effects of the Fabs on elastase release (Figures 9 and 10). Fab concentrations ranged from approximately 20 nM to 300 nM, resulting in Fab:IL-8 molar ratios of 0.2:1 to 3:1. Cytochalasin B (Sigma) was added to the neutrophil suspension at a concentration of 5 µg/ml (using a 5 mg/ml stock solution made up in DMSO), and the cells were incubated for 15 min in a 37°C water bath. Cytochalasin B-treated neutrophils (100 µl) were then added to the IL-8/Fab mixtures. After a 3 hr incubation at room temperature, the neutrophils were pelleted by centrifugation (200 x g for 5 min), and aliquots of the cell-free supernatants were transferred to 96 well plates (30 µl/well). The elastase substrate, methoxysuccinyl-alanyl-prolyl-valyl-p-nitroanilide (Calbiochem, La Jolla, CA), was prepared as a 10 mM stock solution in DMSO and stored at 4°C. Elastase substrate working solution was prepared just prior to use (1.2 mM elastase substrate, 1.2 M NaCl, 0.12 M HEPES, pH 7.2), and 170 µl was added to each sample-containing well. The plates were placed in a 37°C tissue culture incubator for 30 min or until an optical density reading for the positive controls reached at least 1.0. Absorbance was measured at 405 nm using an SLT 340 plate reader (SLT Lab Instruments, Austria).

Figure 9 demonstrates the ability of the chimeric anti-IL-8 Fabs to inhibit elastase release from human neutrophils stimulated by human IL-8; Figure 10 demonstrates the relative abilities of the chimeric anti-IL-8 Fabs to inhibit elastase release from human neutrophils stimulated by rabbit IL-8.

### C. MOLECULAR CLONING OF THE VARIABLE LIGHT AND HEAVY REGIONS OF THE MURINE 5:12.14 (ANTI-IL-8) MONOCLONAL ANTIBODY

Total RNA was isolated from 1 X 10<sup>8</sup> cells (hybridoma cell line ATCC HB-11722) using the procedure described by Chomczynski and Sacchi (Anal. Biochem. 162:156 (1987)). First strand cDNA was synthesized by specifically priming the mRNA with synthetic DNA oligonucleotides designed to hybridize with regions of the murine RNA encoding the constant region of the kappa light chain or the IgG2a heavy chain (the DNA sequence of these regions are published in Sequences of Proteins of Immunological Interest, Kabat, E. A. et al. (1991) NIH Publication 91-3242, V 1-3.). Three primers (SEQ ID NOS: 1-6) were designed for each of the light and heavy chains to increase the chances of primer hybridization and efficiency of first strand cDNA synthesis (Figure 13). Amplification of the first strand cDNA to doublestranded (ds) DNA was accomplished using two sets of synthetic DNA oligonucleotide primers: one forward primer (SEQ ID NOS: 7-9) and one reverse primer (SEQ ID NO: 10) for the light chain variable region amplification (Figure 14) and one forward primer (SEQ ID NOS: 11-14) and one reverse primer (SEQ ID NOS: 15-18) for the heavy chain variable region amplification (Figure 15). The N-terminal sequence of the first eight amino acids of either the light or heavy chains of 5.12.14 was used to generate a putative murine DNA sequence corresponding to this region. (A total of 29 amino acids was sequenced from the N-terminus of both the light chain and heavy chain variable regions using the Edman degradation protein sequencing technique.) This information was used to design the forward amplification primers which were made degenerate in the third position for some codons to increase the chances of primer hybridization to the natural murine DNA codons and also included the unique restriction site, MluI, for both the light chain variable region forward primer and the heavy chain variable region forward primer to

facilitate ligation to the 3' end of the STII element in the cloning vector. The reverse amplification primers were designed to anneal with the murine DNA sequence corresponding to a portion of the constant region of the light or heavy chains near the variable/constant junction. The light chain variable region reverse primer contained a unique BstBI restriction site and the heavy chain variable region reverse primer contained a unique Apal restriction site for ligation to the 5' end of either the human lgG1 constant light or lgG1 constant heavy regions in the vectors, pB13.1 (light chain) and pB14 (heavy chain). The polymerase chain reaction using these primer sets yielded DNA fragments of approximately 400 bp. The cDNA encoding the 5.12.14 light chain variable region was cloned into the vector pB13.1, to form pA51214VL and the 5.12.14 heavy chain variable region was cloned into the vector, pB14, to form pA51214VH. The cDNA inserts were characterized by DNA sequencing and are presented in the DNA sequence (SEQ ID NO: 19) and amino acid sequence (SEQ ID NO: 20) of Figure 16 (murine light chain variable region) and in the DNA sequence (SEQ ID NO: 21) and amino acid (SEQ ID NO: 22) of Figure 17 (murine heavy chain variable region).

#### D. CONSTRUCTION OF A 5.12.14 FAB VECTOR

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In the initial construct, pA51214VL, the amino acids between the end of the 5.12.14 murine light chain variable sequence and the unique cloning site, BstBl, in the human IgG1 constant light sequence were of murine origin corresponding to the first 13 amino acids of the murine IgG1 constant region (Figure 16). Therefore, this plasmid contained a superfluous portion of the murine constant region separating the 5.12.14 murine light chain variable region and the human light chain IgG1 constant region. This intervening sequence would alter the amino acid sequence of the chimera and most likely produce an incorrectly folded Fab. This problem was addressed by immediately truncating the cDNA clone after A109 and re-positioning the BstBI site to the variable/constant junction by the polymerase chain reaction. Figure 18 shows the amplification primers used to make these modifications. The forward primer, VL.front (SEQ ID NO: 23), was designed to match the last five amino acids of the STII signal sequence, including the Mlul cloning site, and the first 4 amino acids of the 5.12.14 murine light chain variable sequence. The sequence was altered from the original cDNA in the third position of the first two codons D1 (T to C) and 12 (C to T) to create a unique EcoRV cloning site which was used for later constructions. The reverse primer, VL.rear (SEQ ID NO: 24), was designed to match the first three amino acids of the human IgG1 constant light sequence and the last seven amino acids of the 5.12.14 light chain variable sequence which included a unique BstBI cloning site. In the process of adding the BstBI site, the nucleotide sequence encoding several amino acids were altered: L106 (TTG to CTT), K107 (AAA to CGA) resulting in a conservative amino acid substitution to arginine, and R108 (CGG to AGA). The PCR product encoding the modified 5.12.14 light chain variable sequence was then subcloned into pB13.1 in a two-part ligation. The MluI-BstBI digested 5.12.14 PCR product encoding the light chain variable region was ligated into Mlul-BstBI digested vector to form the plasmid, pA51214VL'. The modified cDNA was characterized by DNA sequencing. The coding sequence for the 5.12.14 light chain is shown in Figure 19.

Likewise, the DNA sequence between the end of the heavy chain variable region and the unique

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cloning site, ApaI, in the human IgG1 heavy chain constant domain of pA51214VH was reconstructed to change the amino acids in this area from murine to human. This was done by the polymerase chain reaction. Amplification of the murine 5.12.14 heavy chain variable sequence was accomplished using the primers shown in Figure 18. The forward PCR primer (SEQ ID NO: 25) was designed to match nucleotides 867-887 in pA51214VH upstream of the STII signal sequence and the putative cDNA sequence encoding the heavy chain variable region and included the unique cloning site SpeI. The reverse PCR primer (SEQ ID NO: 26) was designed to match the last four amino acids of the 5.12.14 heavy chain variable sequence and the first six amino acids corresponding to the human IgG1 heavy constant sequence which also included the unique cloning site, ApaI. The PCR product encoding the modified 5.12.14 heavy chain variable sequence was then subcloned to the expression plasmid, pMHM24.2.28 in a two-part ligation. The vector was digested with SpeI-ApaI and the SpeI-ApaI digested 5.12.14 PCR product encoding the heavy chain variable region was ligated into it to form the plasmid, pA51214VH'. The modified cDNA was characterized by DNA sequencing. The coding sequence (SEQ ID NO: 29) and amino acid sequence (SEQ ID NO: 30) of Figures 20A-20B.

The first expression plasmid, pantilL-8.1, encoding the chimeric Fab of 5.12.14 was made by digesting pA51214VH' with EcoRV and Bpu11021 to replace the EcoRV-Bpu11021 fragment with a EcoRV-Bpu11021 fragment encoding the murine 5.12.14 light chain variable region of pA51214VL'. The resultant plasmid thus contained the murine-human variable/constant regions of both the light and heavy chains of 5.12.14.

Preliminary analysis of Fab expression using pantilL-8.1 showed that the light and heavy chains were produced intracellularly but very little was being secreted into the periplasmic space of  $\underline{E.~coli}$ . To correct this problem, a second expression plasmid was constructed.

The second expression plasmid, pantilL-8.2, was constructed using the plasmid, pmy187, as the vector. Plasmid pantilL-8.2 was made by digesting pmy187 with Mlul and Sphl and the Mlul (partial)-Sphl fragment encoding the murine 5.12.14 murine-human chimeric Fab of pantilL-8.1 was ligated into it. The resultant plasmid thus contained the murine-human variable/constant regions of both the light and heavy chains of 5.12.14.

The plasmid pantilL-8.2 was deposited on February 10, 1995 with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, U.S.A. (ATCC) and assigned ATTC Accession No. ATCC 97056.

### E. MOLECULAR CLONING OF THE VARIABLE LIGHT AND HEAVY REGIONS OF THE MURINE 6G4.2.5 MONOCLONAL ANTIBODY

Total RNA was isolated from 1x10<sup>8</sup> cells (hybridoma cell line 6G4.2.5) using the procedure described by Chomczynski and Sacchi (Anal. Biochem. 162:156 (1987)). First strand cDNA was synthesized by specifically priming the mRNA with synthetic DNA oligonucleotides designed to hybridize with regions of the murine RNA encoding the constant region of the kappa light chain or the IgG2a heavy chain (the DNA sequence of these regions are published in Sequences of Proteins of Immunological Interest,

Kabat et al. (1991) NIH Publication 91-3242, V 1-3). Three primers (SEQ ID NOS: 31-36) were designed for each the light and heavy chains to increase the chances of primer hybridization and efficiency of first strand cDNA synthesis (Figure 21). Amplification of the first strand cDNA to double-stranded (ds) DNA was accomplished using two sets of synthetic DNA oligonucleotide primers: one forward primer (SEO ID NOS: 37-39) and one reverse primer (SEQ ID NO: 40) for the light chain variable region amplification (Figure 22) and one forward primer (SEQ ID NOS: 41-42) and one reverse primer (SEQ ID NOS: 43-46) for the heavy chain variable region amplification (Figure 23). The N-terminal sequence of the first eight amino acids of either the light or heavy chains of 6G4.2.5 was used to generate a putative murine DNA sequence corresponding to this region. (A total of 29 amino acids were sequenced from the N-terminus of both the light chain and heavy chain variable regions using the Edman degradation protein sequencing technique.) This information was used to design the forward amplification primers which were made degenerate in the third position for some codons to increase the chances of primer hybridization to the natural murine DNA codons and also included the unique restriction site, NsiI, for the light chain variable region forward primer and the unique restriction site, MluI, for the heavy chain variable region forward primer to facilitate ligation to the 3' end of the STII element in the vector, pchimFab. The reverse amplification primers were designed to anneal with the murine DNA sequence corresponding to a portion of the constant region of the light or heavy chains near the variable/constant junction. The light chain variable region reverse primer contained a unique MunI restriction site and the heavy chain variable region reverse primer contained a unique Apal restriction site for ligation to the 5' end of either the human lgG1 constant light or IgG1 constant heavy regions in the vector, pchimFab. The polymerase chain reaction using these primer sets yielded DNA fragments of approximately 400 bp and were cloned individually into the vector, pchimFab, to form p6G425VL and p6G425VH. The cDNA inserts were characterized by DNA sequencing and are presented in the DNA sequence (SEQ ID NO: 47) and amino acid sequence (SEQ ID NO: 48) of Figure 24 (murine light chain variable region) and the DNA sequence (SEQ ID NO: 49) and amino acid sequence (SEQ ID NO: 50) of Figure 25 (murine heavy chain variable region).

#### F. CONSTRUCTION OF A 6G4.2.5 CHIMERIC FAB VECTOR

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In the initial construct, p6G425VL, the amino acids between the end of the 6G4.2.5 murine light chain variable sequence and the unique cloning site. Mun1, in the human IgG1 constant light sequence were of murine origin. These amino acids must match the human IgG1 amino acid sequence to allow proper folding of the chimeric Fab. Two murine amino acids, D115 and S121, differed dramatically from the amino acids found in the loops of the β-strands of the human IgG1 constant domain and were converted to the proper human amino acid residues, V115 and F121, by site-directed mutagenesis using the primers (SEQ ID NOS: 51-54) shown in Figure 26. These specific mutations were confirmed by DNA sequencing and the modified plasmid named p6G425VL'. The coding sequence is shown in the DNA sequence (SEQ ID NO: 55) and amino acid sequence (SEQ ID NO: 56) of Figures 27A-27B.

Likewise, the DNA sequence between the end of the heavy chain variable region and the unique cloning site, Apal, in the human IgG1 heavy chain constant domain of p6G425VH was reconstructed to

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change the amino acids in this area fr m murine to human. This process was facilitated by the discovery of a BstEll site near the end of the heavy chain variable regi n. This site and the Apal site were used for the addition of a synthetic piece of DNA encoding the corresponding IgG human amino acid sequence. The synthetic oligo-nucleotides shown in Figure 26 were designed as complements of one another to allow the formation of a 27 bp piece of ds DNA. The construction was performed as a three-part ligation because the plasmid, p6G425VH, contained an additional BstEll site within the vector sequence. A 5309 bp fragment of p6G425VH digested with Mlul-Apal was ligated to a 388 bp fragment carrying the 6G4.2.5 heavy chain variable region and a 27 bp synthetic DNA fragment encoding the first six amino acids of the human IgG1 constant region to form the plasmid, p6G425VH. The insertion of the synthetic piece of DNA was confirmed by DNA sequencing. The coding sequence is shown in the DNA sequence (SEQ ID NO: 57) and amino acid sequence (SEQ ID NO: 58) of Figures 28A-28B.

The expression plasmid, p6G425chim2, encoding the chimeric Fab of 6G4.2.5 was made by digesting p6G425chimVL' with Mlul and Apal to remove the STII-murine HPC4 heavy chain variable region and replacing it with the Mlul-Apal fragment encoding the STII-murine 6G4.2.5 heavy chain variable region of p6G425chimVH'. The resultant plasmid thus contained the murine-human variable/constant regions of both the light and heavy chains of 6G4.2.5.

The plasmid p6G425chim2 was deposited on February 10, 1995 with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, U.S.A. (ATCC) and assigned ATTC Accession No. 97055.

### 20 G. CONSTRUCTION OF HUMANIZED VERSIONS OF ANTI-IL-8 ANTIBODY 6G4.2.5

The murine cDNA sequence information obtained from the hybridoma cell line, 6G4.2.5, was used to construct recombinant humanized variants of the murine anti-IL-8 antibody. The first humanized variant, F(ab)-1, was made by grafting synthetic DNA oligonucleotide primers encoding the murine CDRs of the heavy and light chains onto a phagemid vector, pEMX1 (Werther et al., J. Immunol, 157: 4986-4995 (1996)), which contains a human 6-subgroup I light chain and a human IgG1 subgroup III heavy chain (Fig. 29). Amino acids comprising the framework of the antibody that were potentially important for maintaining the conformations necessary for high affinity binding to IL-8 by the complementarity-determining regions (CDR) were identified by comparing molecular models of the murine and humanized 6G4.2.5 (F(ab)-1) variable domains using methods described by Carter et al., PNAS 89:4285 (1992) and Eigenbrot, et. al., J. Mol. Biol. 229:969 (1993). Additional humanized framework variants (F(ab) 2-9) were constructed from the information obtained from these models and are presented in Table 4 below. In these variants, the sitedirected mutagenesis methods of Kunkel, Proc. Natl. Acad. Sci USA), 82:488 (1985) were utilized to exchange specific human framework residues with their corresponding 6G4.2.5 murine counterparts. Subsequently, the entire coding sequence of each variant was confirmed by DNA sequencing. Expression and purification of each F(ab) variant was performed as previously described by Werther et. al., supra, with the exception that hen egg white lysozyme was omitted from the purification protocol. The variant antibodies were analyzed by SDS-PAGE, electrospray mass spectroscopy and amino acid analysis.

#### Table 4 - Humanized 6G425 Variants

IC50°

Variant	Version	Template	Changes <sup>a</sup>	Purposeb	Mean	S.D.	N
F(ab)-1	version 1		CDR Swap		63.0	12.3	4
F(ab)-2	version 2	F(ab)-1	PheH67 <i>Ala</i>	packaging w/ CDR H2	106.0	17.0	2
F(ab)-3	version 3	F(ab)-1	ArgH71 <i>Val</i>	packaging w/ CDRs H1, H2	79.8	42.2	4
F(ab)-4	version 6	F(ab)-1	lleH69 <i>Leu</i>	packaging w/ CDR H2	44.7	9.0	3
F(ab)-5	version 7	F(ab)-1	LeuH78 <i>Alu</i>	packaging w/ CDRs H1, H2	52.7	31.0	9
F(ab)-6	version 8	F(ab)-1	IleH69 <i>Leu</i> LeuH78 <i>Ala</i>	combine F(ab)- 4 and -5	34.6	6.7	7
F(ab)-7	version 16	F(ab)-6	LeuH80 <i>Val</i>	packaging w/ CDR H1	38.4	9.1	2
F(ab)-8	version 19	F(ab)-6	ArgH38Lys	packaging w/ CDR H2	14.0	5.7	2
F(ab)-9	version 11	F(ab)-6	GluH6 <i>GIn</i>	packaging w/ CDR H3	19.0	5.1	7
Chimeric <sup>d</sup> F(ab)					11.4	7.0	13
rhu4D5° F(ab)	,	·			>200µM		5

- Amino acid changes made relative to the template used. Murine residues are in bold italics and residue numbering is according to Kabat et al.
  - b Purpose for making changes based upon interactions observed in molecular models of the humanized and murine variable domains.
  - c nM concentration of variant necessary to inhibit binding of iodinated IL-8 to human neutrophils in the competitive binding assay.

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d Chimeric F(ab) is a (F(ab) which carries the murine heavy and light chain variable domains fused to the human light chain kI constant domain and the human heavy chain subgroup III constant domain I respectively.

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e. rhu4D5F(ab) is of the same isotype as the humanized 6G425 F(ab)s and is a humanized anti-HER2 F(ab) and therefore should not bind to IL8.

The first humanized variant, F(ab)-1, was an unaltered CDR swap in which all the murine CDR amino acids defined by both x-ray crystallography and sequence hypervariability were transferred to the human framework. When the purified F(ab) was tested for its ability to inhibit 125 I-IL-8 binding to human neutrophils according to the methods described in Section (B)(1) above, a 5.5 fold reduction in binding affinity was evident as shown in Table 4 above. Subsequent versions of F(ab)-1 were engineered to fashion the 3-dimensional structure of the CDR loops into a more favorable conformation for binding IL-8. The relative affinities of the F(ab) variants determined from competition binding experiments using human neutrophils as described in Section (B)(1) above are presented in Table 4 above. A slight decrease in IL-8 binding (<2 fold) was observed for F(ab)-2-3 while only slight increases in IL-8 binding were noted for F(ab)3-5. Variant F(ab)-6 had the highest increase in affinity for IL-8 (approximately 2 fold), exhibiting an IL-8 binding affinity of 34.6nM compared to the F(ab)-1 IL-8 binding affinity of 63nM. The substitutions of murine Leu for Ile at H69 and murine Ala for Leu at H78 are predicted to influence the packing of CDRs H1 and H2. Further framework substitutions using the F(ab)-6 variant as template were made to bring the binding affinity closer to that of the chimeric F(ab). In-vitro binding experiments revealed no change in affinity for F(ab)-7 (38.4nM) but a significant improvement in affinity for F(ab)-8/9 of 14nM and 19 nM, respectively. By analysis of a 3-D computer-generated model of the anti-IL-8 antibody, it was hypothesized that the substitution of murine Lys for Arg at H38 in F(ab)-8 influences CDR-H2 while a change at H6 of murine Gln for Glu in F(ab)-9 affects CDR-H3. Examination of the human antibody sequences with respect to amino acid variability revealed that the frequency of Arg at residue H38 is >99% whereas residue H6 is either Gln ~20% or Glu ~80% (Kabat et. al., Sequences of Proteins of Immunological Interest 5th Ed. (1991)). Therefore, to reduce the likelihood of causing an immune response to the antibody, F(ab)-9 was chosen over F(ab)-8 for further affinity maturation studies. Variant F(ab)-9 was also tested for its ability to inhibit IL-8-mediated chemotaxis (Fig. 30). This antibody was able to block neutrophil migration induced by wild-type human IL-8, human monomeric IL-8 and Rhesus IL-8 with IC<sub>50</sub>=s of approximately 12nM, 15nM, and 22nM, respectively, in IL-8 mediated neutrophil chemotaxis inhibition assays performed as described in Section (B)(2) above. The amino acid sequence for variant F(ab)-8 is provided in Fig. 31c. The F(ab)-8 was found to block human and rhesus IL-8-mediated chemotaxis with IC50=s of 12nM and 10nM, respectively, in 1L-8 mediated neutrophil chemotaxis inhibition assays performed as described in Section (B)(2) above.

### H. <u>CONSTRUCTION OF AN ANTI-IL-8-GENE III FUSION PROTEIN FOR PHAGE DISPLAY AND ALANINE SCANNING MUTAGENESIS</u>

An expression plasmid, pPh6G4.V11, encoding a fusion protein (heavy chain of the humanized 6G4.2.5 version 11 antibody and the M13 phage gene-III coat protein) and the light chain of the humanized 6G4.2.5 version 11 antibody was assembled to produce a monovalent display of the anti-IL-8 antibody on

phage particles. The construct was made by digesting the plasmid, pFPHX, with EcoRV and Apal to remove the existing irrelevant antibody coding sequence and replacing it with a 1305bp EcoRV-Apal fragment from the plasmid, p6G4.V11, encoding the humanized 6G4.2.5 version 11 anti-IL-8 antibody. The translated sequence of the humanized 6G4.2.5 version 11 heavy chain (SEQ ID NO: 66), peptide linker and gene III coat protein (SEQ ID NO: 67) is shown in Fig. 31A. The pFPHX plasmid is a derivative of phGHam-3 which contains an in-frame amber codon (TAG) between the human growth hormone and gene-III DNA coding sequences. When transformed into an amber suppressor strain of E. coli, the codon (TAG) is read as Glutamate producing a growth hormone (hGH)-gene III fusion protein. Likewise, in a normal strain of E. coli, the codon (TAG) is read as a stop preventing translational read-through into the gene-III sequence and thus allowing the production of soluble hGH. The pGHam-3 plasmid is described in Methods: A Companion to Methods in Enzymology, 3:205 (1991). The final product, pPh6G4.V11, was used as the template for the alanine scanning mutagenesis of the CDRs and for the construction of randomized CDR libraries of the humanized 6G4.V11 antibody.

#### I. ALANINE SCANNING MUTAGENESIS OF HUMANIZED ANTIBODY 6G4.2.5 VERSION 11

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The solvent exposed amino acid residues in the CDRs of the humanized anti-IL-8 6G4.2.5 version 11 antibody (h6G4V11) were identified by analysis of a 3-D computer-generated model of the anti-IL-8 antibody. In order to determine which solvent exposed amino acids in the CDRs affect binding to interleukin-8, each of the solvent exposed amino acids was individually changed to alanine, creating a panel of mutant antibodies wherein each mutant contained an alanine substitution at a single solvent exposed residue. The alanine scanning mutagenesis was performed as described by Leong et. al., J. Biol. Chem., 269: 19343 (1994)).

The IC<sub>50</sub>'s (relative affinities) of h6G4V11 wt and mutated antibodies were established using a Competition Phage ELISA Assay described by Cunningham et. al., (EMBO J. 13:2508 (1994)) and Lee et. al., (Science 270:1657 (1995)). The assay measures the ability of each antibody to bind IL-8 coated onto a 96-well plate in the presence of various concentrations of free IL-8 (0.2 to 1uM) in solution. The first step of the assay requires that the concentrations of the phage carrying the wild type and mutated antibodies be normalized, allowing a comparison of the relative affinities of each antibody. The normalization was accomplished by titering the phage on the IL-8 coated plates and establishing their EC50. Sulfhydryl coated 96-well binding plates (Corning-Costar; Wilmington, MA) were incubated with a 0.1mg/ml solution of K64C IL-8 (Lysine 64 is substituted with Cysteine to allow the formation of a disulfide bond between the free thiol group of K64C IL-8 and the sulfhydryl coated plate, which results in the positioning of the IL-8 receptor binding domains towards the solution interface) in phosphate buffered saline (PBS) pH 6.5 containing 1mM EDTA for 1 hour at 25EC followed by three washes with PBS and a final incubation with a solution of PBS containing 1.75mg/ml of L-cysteine-HCl and 0.1M NaHCO, to block any free reactive sulfhydryl groups on the plate. The plates were washed once more and stored covered at 4EC with 200ul of PBS/well. Phage displaying either the reference antibody, h6G4V11, or the mutant h6G4V11 antibodies were grown and harvested by PEG precipitation. The phage were resuspended in 500ul 10mM Tris-HCl pH

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7.5, 1mM EDTA and 100mM NaCl and held at 4EC for no longer than 3 hours. An aliquot of each phage was diluted 4-fold in PBS containing 0.05% Tween-20 (BioRad, Richmond, Ca.) and 0.5% BSA RIA grade (Sigma, St. Louis, Mo.) (PBB) and added to IL-8 coated plates blocked for at least 2 hours at 25EC with 50mg/ml skim milk powder in 25mM Carbonate Buffer pH 9.6. The phage were next serially diluted in 3 fold steps down the plate from well A through H. The plates were incubated for 1 hour at 25EC followed by nine quick washes with PBS containing 0.05% Tween-20 (PBST). The plates were then incubated with a 1:3200 dilution of rabbit anti-phage antibody and a 1:1600 dilution of secondary goat-anti-rabbit Fc HRPconjugated antibody for 15 minutes at 25EC followed by nine quick washes with PBST. The plates were developed with 80ul/well of 1mg/ml OPD (Sigma, St. Louis, Mo) in Citrate Phosphate buffer pH 5.0 containing 0.015% H<sub>2</sub>O<sub>2</sub> for 4 minutes at 25EC and the reaction stopped with the addition of 40ul of 4.5M H<sub>2</sub>SO<sub>4</sub>. The plates were analyzed at wavelength 8<sub>492</sub> in a SLT model 340ATTC plate reader (SLT Lab The individual EC<sub>50</sub>=s were determined by analyzing the data using the program Instruments). Kaleidagraph (Synergy Software, Reading, Pa.) and a 4-parameter fit equation. The phage held at 4EC were then immediately diluted in PBB to achieve a final concentration corresponding to their respective EC50 or target OD492 for the competition segment of the experiment, and dispensed into a 96 well plate containing 4-fold serial dilutions of soluble IL-8 ranging from 1uM in well A and ending with 0.2uM in well H. Using a 12-channel pipet, 100ul of the phage/IL-8 mixture was transferred to an IL-8 coated 96-well plate and executed as described above. Each sample was done in triplicate - 3 columns/sample.

Table 5 - Relative Affinities (IC50) for Alanine-scan Anti-IL-8 6G4V11 CDR Mutants

CDR	Amino Acid Residue	Avg IC50 (nM)	Std Dev
VII	Reference	11.5	6.4
CDR-L1	S26	6.3	2.9
	Q27	10.2	2.4
	S28	14.2	5.2
	V30	29.1	12.3
	H31	580.3	243.0
	133	64.2	14.6
	N35	3.3	0.7
	T36	138.0	nd
	Y37	NDB	nd
CDR-L2	K55	24.2	14.9
	V56	15.5	3.8
	\$57	12.4	4.0
	N58	17.6	3.7
	R59	nd .	nd
CDR-L3	S96	10.8	4.4
<del></del>	T97	70.6	55.2

·CDR	Amino Acid Residue	Avg IC50 (nM)	Std Dev
	H98	8.0	1.2
	V99	19.6	1.9
CDR-H1	S28	8.6	3.1
	S30	nd	nd
	S31	7.8	2.5
	H32	13.3	5.8
	Y53	48.2	15.8
CDR-H2	Y50	35.6	13.0
	D52	13.3	7.5
<del></del>	S53	6.0	3.4
	N54	96.0	5.8
	E56	15.8	4.5
	T57	8.4	1.6
	T58	11.3	1.8
	Y59	9.1	3.7
	Q61	12.6	6.4
	K64	18.5	12.1
CDR-H3	D96	NDB	nd
	Y97	NDB	nd
<del></del>	R98	36.6	15.3
	Y99 #	199.5	nd
	N100	278.3	169.4
	D102	159.2	44
	W103	NDB	nd
	F104	NDB	nd
	F105	209.4	72.3
	D106	25.3	21.7

Each sample performed in triplicate/experiment.

NDB = No Detectable Binding /nd = value not determined\*

Residue numbering is according to Kabat et al.

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The results of the alanine-scan are summarized in Table 5 above. The alanine substitutions in of many of the mutant antibodies had little or no adverse effects (<3 fold) on the binding affinity for IL-8. Mutants that were found to exhibit no detectable binding of IL-8 (NDB) presumably contained disruptions in the conformational structure of the antibody conferred by crucial structural or buried amino acids in the CDR. Based on the results of the scan, CDR-H3 (heavy chain, 3rd CDR) was identified as the dominant binding epitope for binding IL-8. Alanine substitutions in this CDR resulted in a 3 to >26 fold decrease in binding affinity. The amino acids, Y597, Y599 and D602 are of particular interest because it was determined from the computer generated model of the anti-IL-8 antibody that these residues are solvent exposed and that these residues might participate in hydrogen bonding or charge interactions with IL-8 or other amino acids of the antibody that influence either binding to IL-8 r the conformation of the CDR-H3

loop structure. (See the model depicted in Fig. 32). Unexpected increases in binding affinity (1.8 > 2.7 fold) were noted for S528 and S531 of CDR-H1 and S553 of CDR-H2.

Surprisingly, a significant increase in binding affinity was observed in the alanine mutant N35A located in CDR-L1 (light chain, 1st CDR). A 3-6 fold increase in affinity was observed compared to the wild-type h6G4V11 antibody. This augmentation of IL-8 binding could be the result of the close proximity of N35A to CDR-H3. The alanine substitution may have imparted a slight change in the conformation of CDR-L1 which alters the packing interaction of neighboring amino acid residues on CDR-H3, thereby tweaking the loop of CDR-H3 into a conformation that facilitates more appropriate contacts with IL-8. Similarly, N35A may also influence the orientation of amino acids in CDR-L1 or its interaction directly with IL-8. Unexpected increases in affinity (~2 fold) were also observed for S26 of CDR-L1 and H98 of CDR-L3.

#### J. CHARACTERIZATION OF HUMANIZED ANTI-IL-8 ANTIBODY 6G4V11N35A

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Soluble 6G4V11N35A Fab antibody was made by transforming an amber non-suppressor strain of E. coli, 34B8, with pPh6G4.V11 and growing the culture in low phosphate medium for 24 hours. The periplasmic fraction was collected and passed over a Hi-Trap Protein-G column (Pharmacia, Piscataway, NJ.) followed by a desalting and concentration step. The protein was analyzed by SDS-PAGE, mass spectrometry and amino acid analysis. The protein had the correct size and amino acid composition (Fig. 35). The 6G4V11N35A Fab was tested for its ability to inhibit 125 I-IL-8 binding to human neutrophils and to inhibit IL-8 mediated neutrophil chemotaxis as described in Section (B)(1) and (B)(2) above. As shown in Fig. 33, hybridoma-derived intact murine antibody (6G4 murine mAB), recombinant 6G4 murine-human chimera Fab, recombinant humanized Fab versions 1 and 11, and 6G4V11N35A Fab were found to inhibit 125<sub>1-1L-8</sub> binding to human neutrophils with an average IC<sub>50</sub> of 5nM, 8nM, 40nM, 10nM and 3nM, respectively. The 6G4V11N35A Fab had at least a 2-fold higher affinity than the 6G4.2.5 chimera Fab and a 3-fold higher affinity than 6G4V11. As shown in Fig. 34, the 6G4V11N35A Fab was found to inhibit IL-8 mediated neutrophil chemotaxis induced by both wild type and monomeric human IL-8, and by two different animal species of IL-8, namely, rabbit and rhesus. The irrelevant isotype control Fab (4D5) did not inhibit neutrophil migration. The average IC50 values were 3nM (wt IL-8), 1 nM (monomeric IL-8), 5nM (Rabbit IL-8), and 10nM (Rhesus IL-8).

#### K. CONSTRUCTION OF A 6G4V11N35A F(ab'), LEUCINE ZIPPER

Production of a F(ab')<sub>2</sub> version of the humanized anti-IL-8 6G4V11N35A Fab was accomplished by constructing a fusion protein with the yeast GCN4 leucine zipper. The expression plasmid p6G4V11N35A.F(ab')<sub>2</sub> was made by digesting the plasmid p6G425chim2.fab2 with the restriction enzymes bsal and apal to remove the DNA sequence encoding the 6G4.2.5 murine-human chimeric Fab and replacing it with a 2620bp bsal-apal fragment from pPh6G4.V11N35A. The plasmid p6G425chim2.fab2 is a derivative of pS1130 which encodes a fusion protein (the GCN4 leucine zipper fused to the heavy chain of

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anti-CD18) and the light chain of anti-CD18 antibody. The expression plasmid p6G4V11N35A.F(ab')<sub>2</sub> was deposited on February 20, 1996 with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, U.S.A. (ATCC) and assigned ATCC Accession No. 97890. A pepsin cleavage site in the hinge region of the antibody facilitates the removal of the leucine zipper leaving the two immunoglobin monomers joined by the cysteines that generate the interchain disulfide bonds. The DNA and protein sequence of the h6G4V11N35A.F(ab')<sub>2</sub> are depicted in Figs. 35-37.

An expression host cell was obtained by transforming E. coli strain 49D6 with p6G4V11N35A.F(ab')<sub>2</sub> essentially as described in Section (II)(3)(C) above. The transformed host E. coli 49D6 (p6G4V11N35A.F(ab')<sub>2</sub>) was deposited on February 20, 1997 at the ATCC and assigned ATCC Accession No. 98332. Transformed host cells were grown in culture, and the 6G4V11N35A F(ab')<sub>2</sub> product was harvested from the host cell periplasmic space essentially as described in Section (II)(3)(F) above.

#### L. CHARACTERIZATION OF THE HUMANIZED 6G4V11N35A F(ab'), LEUCINE ZIPPER

The 6G4V11N35A Fab and  $F(ab')_2$  were tested for their ability to inhibit <sup>125</sup>I-IL-8 binding to neutrophils according to the procedures described in Section (B)(1) above. The displacement curves from a representative binding experiment performed in duplicate is depicted in Fig. 38. Scatchard analysis of this data shows that 6G4V11N35A  $F(ab')_2$  inhibited <sup>125</sup>I-IL-8 binding to human neutrophils with an average IC<sub>50</sub> of 0.7 nM (+/- 0.2). This is at least a 7 fold increase in affinity compared to the hybridoma-derived intact murine antibody (average IC<sub>50</sub> of 5 nM) and at least a 2.8 fold increase in affinity over the Fab version (average IC<sub>50</sub> of 2 nM).

The 6G4V11N35A F(ab')2 was also tested for its ability to inhibit IL-8 mediated neutrophil chemotaxis according to the procedures described in Section (B)(2) above. The results of a representative chemotaxis experiment performed in quadruplicate are depicted in Fig. 39. As shown in Fig. 39, the 6G4V11N35A F(ab')<sub>2</sub> inhibited human IL-8 mediated neutrophil chemotaxis. The 6G4V11N35A F(ab')<sub>2</sub> exhibited an average IC50 value of 1.5nM versus 2.7nM for the 6G4V11N35A Fab, which represents an approximately 2 fold improvement in the antibody's ability to neutralize the effects of IL-8. The irrelevant isotype control Fab (4D5) did not inhibit neutrophil migration. Furthermore, the 6G4V11N35A F(ab')2 antibody retained its ability to inhibit IL-8 mediated neutrophil chemotaxis by monomeric IL-8 and by two different animal species of IL-8, namely rabbit and rhesus, in neutrophil chemotaxis experiments conducted as described above. An individual experiment is shown in Fig. 40. The average IC50 values were 1nM IL-8). (Rhesus 2.0nM (monomeric IL-8), 4nM (Rabbit IL-8), and

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### M. RANDOM MUTAGENESIS OF LIGHT CHAIN AMINO ACID (N35A) IN CDR-L1 OF HUMANIZED ANTIBODY 6G4V11

A 3-fold improvement in the IC<sub>50</sub> for inhibiting <sup>125</sup>I-IL-8 binding to human neutrophils was observed when alanine was substituted for asparagine at position 35 in CDR-L1 (light chain) of the humanized 6G4V11 mAb as described in Section (I) above. This result might be attributed to an improvement in the contact between the antigen-antibody binding interfaces as a consequence of the replacement of a less bulky nonpolar side chain (R-group) that may have altered the conformation of CDR-L1 or neighboring CDR-H3 (heavy chain) to become more accessible for antigen docking. The acceptance of alanine at position 35 of CDR-L1 suggested that this position contributed to improved affinity and that an assessment of the re-modeling of CDR loops / antigen-binding region(s) by other amino acids at this location was warranted. Selection of an affinity matured version of the humanized 6G4.V11 mAB (Kunkel, T. A., <u>Proc. Natl. Acad. Sci. USA</u>, 82:488 (1995)) was accomplished by randomly mutagenizing position 35 of CDR-L1 and constructing an antibody-phage library. The codon for Asparagine (N) at position 35 of CDR-L1, was targeted for randomization to any of the 20 known amino acids.

Initially, a stop template, pPh6G4.V11-stop, was made to eliminate contaminating wild-type N35 sequence from the library. This was accomplished by performing site-directed mutagenesis (Muta-Gene Kit, Biorad, Ricmond, CA) of pPH6G4V11 (described in Section (H) above) to replace the codon (AAC) for N35 with a stop codon (TAA) using the primer SL.97.2 (SEQ ID NO: )(Figure 42). The incorporation of the stop codon was confirmed by DNA sequencing. Subsequently, uracil containing single-stranded DNA derived from E. coli CJ236 transformed with the stop template was used to generate an antibodyphage library following the method described by Lowman (Methods in Molecular Biology, 87 Chapter 25: 1-15 (1997). The variants generated from this library were predicted to produce a collection of antibodies containing one of the 20 known amino acids at position N35 in CDR-L1. The amino acid substitutions were accomplished by site-directed mutagenesis using the degenerate oligonucleotide primer (SL.97.3) with the sequence NNS (N = A/G/T/C.; S = G/C; ) (SEQ ID NO: )(Figure 42). This codon usage should allow for the expression of any of the 20 amino acids - including the amber stop codon (TAG). The collection of antibody-phage variants was transfected into E. coli strain XL-1 blue (Stratagene, San Diego, CA) by electroporation and grown at 37°C overnight to amplify the library. Selection of tight binding humanized 6G4V11 Fab's were accomplished by panning the library on IL-8 coated 96-well plates as described in Section (I) above. Prior to panning, the number of phage/library was normalized to 1.1x10<sup>13</sup> phage/ml (which produces a maximum OD<sub>270</sub> reading = 1 OD unit) and IL-8 coated plates were incubated with blocking solution (25mN Carbonate buffer containing 50mg/ml skim milk) for 2 hours before the addition of phage (each sort used eight IL-8 coated wells/library). After the blocking and washing steps, every sort began with the addition of 100ul of antibody-phage (titered at 1.1x1013 phage/ml) to each of eight IL-8 coated wells followed by an 1 hour incubation at 25°C. The nonspecifically bound antibody-phage were removed by 10 quick washes with PBS-0.05% Tween 20 (PBS-

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Tween). For sort #1, a low stringency wash (100ul PBS-Tween/well for 10 minutes at 25°C) was employed to capture the small proportion of tight binding antibody-phage bound to the immobilized IL-8. The antibody-phage variants specifically bound to IL-8 were eluted with 100ul/well of 200mM Glycine pH 2.0 for 5 minutes at 25°C. The eluted antibody-phage variants from the 8 wells were then pooled and neutralized with 1M Tris-HCl pH 8.0 (1/3 the elution volume). The phage were titered and propagated as described in Section (I) above. The stringency of the washes were successively increased with each round of panning depending upon the percent recovery of phage at the end of a sort. The wash conditions were as follows: sort #2 (4 x 15 minute intervals; total time = 60 minutes) and sort #3 (either #3a: 8 x 15 minute intervals or #3b: 12 x 10 minute intervals; total time = 120 minutes). The total number of phage recovered was progressively reduced after each sort suggesting that non- or weak- binders were being selected against. The recovery of the negative control (the antibody-phage stop variant) was constant throughout the panning (approximately 0.0001 to 0.00001 percent).

Eighteen random variants from sort #3 were analyzed by DNA sequencing to look for an amino acid consensus at position 35 of CDR-L1. The data presented in Figure 43A showed that Glycine occupied position 35 in 33% of the variants sequenced. However, after correcting for the number of NNS codon combinations/amino acid, the frequency of Glycine was reduced to 16.6%. Glutamic Acid was represented with the highest frequency (22%) followed by Aspartic Acid and Glycine (16.6%). The frequencies of recovery of the wild-type Asparagine and substituted Alanine were only 5.6%. Interestingly, the high frequency of Glycine may suggest that a much wider range of conformations might be allowed for the loop of CDR-L1 which may be attributed to the reduction in steric hindrance of bond angle (φ-ψ) pairing as a result of the single hydrogen atom as the side chain. Conversely, Glutamic Acid at position 35 might restrict the flexibility of the loop by imposing less freedom of rotation imposed by the more rigid and bulky charged polar side chain.

Soluble Fab's of the affinity matured variants (N35G, N35D, N35E and N35A) were made as described in Section (J) above for evaluating their ability to block IL-8 binding. As shown in Figure 43B, variants N35A, N35D, N35E and N35G were found to inhibit <sup>125</sup>I-IL-8 binding to human neutrophils with an approximate IC<sub>50</sub> of 0.2nM, 0.9nM, 0.1nM and 3.0nM, respectively. All of the affinity matured variants showed an improvement in binding IL-8 ranging from 3 - 100 fold compared to the humanized 6G4V11 mAb. The affinity-matured variant, 6G4V11N35E, was 2-fold more potent in blocking IL-8 binding to human neutrophils than the alanine-scan variant, 6G4V11N35A.

Equilibrium and kinetic measurements of variants 6G4V11N35A and 6G4V11N35E were determined using KinEXA<sup>TM</sup> automated immunoassay system (Sapidyne Instruments Inc., Idaho City, ID) as described by Blake *et al.*, J. Biol. Chem. 271: 27677 (1996). The procedure for preparing the antigencoated particles was modified as follows: 1 ml of activated agarose beads (Reacti-Gel 6X; Pierce, Rockford, IL) were coated with antigen in 50mM Carbonate buffer pH 9.6 containing 20ug/ml of human IL-8 and incubated with gentle agitation on a rocker overnight at 25°C. The IL-8 coated beads were then

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washed twic with 1M Tris-HCl pH 7.5 to inactivate any unreactive groups on the beads and blocked with Superblock (Pierce, Rockford, IL) for 1 hour at 25C to reduce non-specific binding. The beads were resuspended in assay buffer (0.1% bovine serum albumin in PBS) to a final volume of 30 ml. A 550ul aliquot of the IL-8 coated bead suspension was used each time to pack a fresh 4mm high column in the KinEXA observation cell. The amount of unbound antibody from the antibody-antigen mixtures captured by the IL-8-coated beads in both the equilibrium and kinetic experiments was quantified using a fluorescently labeled secondary antibody. Murine 6G4.2.5 was detected with a R-PE AffiniPure F(ab')<sub>2</sub> goat anti-mouse IgG, Fc fragment specific 2° antibody (Jackson Immuno Research Laboratories, West Grove, PA) and humanized affinity matured N35A (Fab and F(ab')<sub>2</sub>) and N35E Fab were detected with a R-PE AffiniPure F(ab')<sub>2</sub> donkey anti-human IgG (H+L) 2° antibody (Jackson Immunoresearch Laboratories, West Grove, PA); both at a 1:1000 dilution.

Equilibrium measurements were determined by incubating a constant amount of anti-IL-8 antibody (0.005ug/ml) with various concentrations of human IL-8 (0, 0.009, 0.019, 0.039, 0.078, 0.156, 0.312, 0.625, 1.25, 2.5nM). The antibody-antigen mixture was incubated for 2 hours at 25°C to allow the molecules to reach equilibrium. Subsequently, each sample was passed over a naive IL-8 coated bead pack in the KinEXA observation cell at a flow rate of 0.5ml/minute for a total of 9 minutes/sample. The equilibrium constant (Kd) was calculated using the software provided by Sapidyne Instruments Inc.

Rates of association (ka) and dissociation (kd) were determined by incubating together a constant amount of antibody and antigen, and measuring the amount of uncomplexed anti-IL-8 bound to the IL-8 coated beads over time. The concentration of antibody used in the kinetic experiments was identical to that used in the equilibrium experiment described above. Generally, the amount of human IL-8 used was the concentration derived from the binding curves of the equilibrium experiment that resulted in 70% inhibition of anti-IL-8 binding to the IL-8 coated beads. Measurements were made every 15 minutes to collect approximately nine data points. The ka was calculated using the software provided by Sapidyne Instruments, Inc. The off rate was determined using the equation: kd = Kd/ka.

Figure 44 shows the equilibrium constants (Kd) for the affinity matured variants 6G4V11N35E and 6G4V11N35A Fab's were approximately 54pM and 114pM, respectively. The improvement in affinity of 6G4V11N35E Fab for IL-8 can be attributed to a 2-fold faster rate of association (K<sub>on</sub>) of 4.7x10<sup>6</sup> for 6G4V11N35E Fab versus 2.0x10<sup>6</sup> for 6G4V11N35A F(ab')<sub>2</sub>. (The Kd of the 6G4V11N35A F(ab')<sub>2</sub> and 6G4V11N35A Fab are similar.) The dissociation rates (K<sub>off</sub>) were not significantly different. Molecular modeling suggests that substitution of Aspargine with Glutamic Acid might either affect the antibody's interaction with IL-8 directly or indirectly by neutralizing the charge of neighboring residues R98 (CDR-H3) or K50 (CDR-L2) in the CDR's to facilitate contact with IL-8. Another effect might be the formation of a more stable loop conformation for CDR-L1 that could have facilitated more appropriate contacts of other CDR-L1 loop residues with IL-8. The DNA (SEQ ID NO: ) and amino acid (SEQ ID NO: )

sequences of p6G4V11N35E.Fab showing the Asparagine to Glutamic Acid substitution in the light chain ar presented in Figure 45.

#### N: CHARACTERIZATION OF HUMANIZED ANTI-IL-8 VARIANT 6G4V11N35E Fab

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The affinity matured Fab variant, 6G4V11N35E, was tested for its ability to inhibit IL-8 mediated neutrophil chemotaxis as described in Section (B)(2) above. The reuseable 96-well chemotaxis chamber described in Section (B)(2) was replaced with endotoxin-free disposable chemotaxis chambers containing 5-micron PVP-free polycarbonate filters (ChemoTx101-5, Neuro Probe, Inc. Cabin John, MD). As illustrated in Figure 46, variant N35E effectively blocks IL-8 mediated neutrophil chemotaxis induced by a 2nM stimulus of either rabbit or human IL-8. In fact, the level of inhibition at antibody concentrations between 3.7nM - 33nM was not significantly different from the buffer control indicating variant N35E could completely inhibit this response. The IC<sub>50</sub>'s for both rabbit and human IL-8 were approximately 2.8nM and 1.2nM, respectively. The irrelevant isotype control Fab (4D5) did not inhibit neutrophil migation indicating the results observed for the affinity matured variant, N35E, is IL-8 specific.

#### O. CONSTRUCTION OF HUMANIZED 6G4V11N35E F(ab')<sub>2</sub> LEUCINE ZIPPER

A F(ab')<sub>2</sub> expression plasmid for 6G4V11N35E was constructed using methods similar to those described in Section (K) above. The expression plasmid, p6G4V11N35E.F(ab')<sub>2</sub>, was made by digesting the plasmid p6G4V11N35A.F(ab')<sub>2</sub> (described in Section (K) above) with the restriction enzymes Apal and Ndel to isolate a 2805 bp fragment encoding the heavy chain constant domain -GCN4 leucine zipper and ligating it to a 3758 bp Apal-Ndel fragment of the pPH6G4V11N35E phage display clone (encoding 6G4V11N35E Fab) obtained as described in Section (M) above. The integrity of the entire coding sequence was confirmed by DNA sequencing.

### P. CONSTRUCTION OF THE FULL LENGTH HUMANIZED 6G4V11N35A IgG EXPRESSION PLASMID

The full length IgG<sub>1</sub> version of the humanized anti-IL8 variant 6G4V11N35A was made using a dicistronic DHFR-Intron expression vector (Lucas et al., Nucleic Acids Res.,24: 1774-1779 (1996)) which contained the full length recombinant murine-human chimera of the 6G4.2.5 anti-IL8 mAb. The expression plasmid encoding the humanized variant 6G4V11N35A was assembled as follows. First an intermediate plasmid (pSL-3) was made to shuttle the sequence encoding the variable heavy chain of humanized anti-IL-8 variant 6G4V11N35A to pRK56G4chim.2Vh - which contains the variable heavy region of the chimeric 6G4.5 anti-IL8 antibody. The vector pRK56G4chim.Vh was digested with PvuII and ApaI to remove the heavy chain variable region of the chimeric antibody and religated with an 80bp PvuII - Xhol synthetic oligonucleotide (encoding Leu4 to Phe29 of 6G4V11N35A) (Fig. 47) and a 291bp Xhol - ApaI fragment from p6G4V11N35A.7 carrying the remainder of the variable heavy chain sequence of 6G4V11N35A to create pSL-3. This intermediate plasmid was used in conjunction with 2 other plasmids, p6G4V11N35A.F(ab')<sub>2</sub> and p6G425chim2.choSD, to create the mammalian expression plasmid,

p6G4V11N35AchoSD.9 (identified as p6G425V11N35A.choSD in a deposit made on December 16, 1997 with the ATCC and assigned ATCC Accession No. 209552). This expression construct was assembled in a 4-part ligation using the following DNA fragments: a 5,203bp ClaI - BlpI fragment encoding the regulatory elements of the mammalian expression plasmid (p6G425 chim2.choSD), a 451bp ClaI - ApaI fragment containing the heavy chain variable region of the humanized 6G4V11N35A antibody (pSL-3), a 1,921bp ApaI - EcoRV fragment carrying the heavy chain constant region of 6G4V11N35A (p6G425chim2.choSD) and a 554bp EcoRV - BlpI fragment encoding the light chain variable and constant regions of 6G4V11N35A (p6G4V11N35A.F(ab')<sub>2</sub>). The DNA sequence (SEQ ID NO: ) of clone p6G4V11N35A.choSD.9 was confirmed by DNA sequencing and is presented in Figure 48.

## Q. CONSTRUCTION OF THE FULL LENGTH HUMANIZED 6G4V11N35E IgG EXPRESSION PLASMID

A mammalian expression vector for the humanized 6G4V11N35E was made by swapping the light chain variable region of 6G4V11N35A with 6G4V11N35E as follows: a 7,566bp EcoRV - Blp1 fragment (void of the 554bp fragment encoding the light chain variable region of 6G4V11N35A) from p6G4V11N35A.choSD.9 was ligated to a 554bp EcoRV - Blp1 fragment (encoding the light chain variable region of 6G4V11N35E) from pPH6G4V11N35E.7. The mutation at position N35 of the light chain of p6G4V11N35E.choSD.10 was confirmed by DNA sequencing.

#### R. STABLE CHO CELL LINES FOR VARIANTS N35A AND N35E

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For stable expression of the final humanized IgG1 variants (6G4V11N35A and 6G4V11N35E), Chinese hamster ovary (CHO) DP-12 cells were transfected with the above-described dicistronic vectors (p6G4V11N35A.choSD.9 and p6G4V11N35E.choSD.10, respectively) designed to coexpress both heavy and light chains (Lucas et al., Nucleic Acid Res. 24:1774-79 (1996)). Plasmids were introduced into CHO DP12 cells via lipofection and selected for growth in GHT-free medium (Chisholm, V. High efficiency gene transfer in mammalian cells. In: Glover, DM, Hames, BD. DNA Cloning 4. Mammalian systems. Oxford Univ. Press, Oxford pp 1-41 (1996)). Approximately 20 unamplified clones were randomly chosen and reseeded into 96 well plates. Relative specific productivity of each colony was monitored using an ELISA to quantitate the full length human IgG accumulated in each well after 3 days and a fluorescent dye, Calcien AM, as a surrogate marker of viable cell number per well. Based on these data, several unamplified clones were chosen for further amplification in the presence of increasing concentrations of methotrexate. Individual clones surviving at 10, 50, and 100 nM methotrexate were chosen and transferred to 96 well plates for productivity screening. One clone for each antibody (clone#1933 alL8.92 NB 28605/12 for 6G4V11N35A; clone#1934 aIL8.42 NB 28605/14 for 6G4V11N35E), which reproducibly exhibited high specific productivity, was expanded in T-flasks and used to inoculate a spinner culture. After several passages, the suspension-adapted cells were used to inoculate production cultures in GHT-containing, serum-free media supplemented with various hormones and protein hydrolysates. Harvested cell culture fluid containing recombinant humanized anti-IL8 was purified using protein A-Sepharose CL-4B. The purity after this step was approximately 99%. Subsequent purification to homogeneity was carried out

using an ion exchange chromatography step. Production titer f the humanized 6G4V11N35E IgG1 antibody after the first round of amplification and 6G4V11N35A IgG1 after the second round of amplification were 250mg/L and 150mg/L, respectively.

#### S. CHARACTERIZATION OF THE HUMANIZED 6G4V11N35A/E IgG VARIANTS

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The humanized full length IgG variants of 6G4.2.5 were tested for their ability to inhibit  $^{125}$ I-IL-8 binding and to neutralize activation of human neutrophils; the procedures are described in Sections (B)(1) and (B)(2) above. As shown in Figure 49, the full length IgG1 forms of variants 6G4V11N35A and 6G4V11N35E equally inhibited  $^{125}$ I-IL-8 binding to human neutrophils with approximate IC<sub>50</sub>'s of 0.3nM and 0.5nM, respectively. This represents a 15 - 25 fold improvement in blocking binding of IL-8 compared to the full length murine mAb (IC<sub>50</sub> = 7.5nM). Similarly, the two anti-IL-8 variants showed equivalent neutralizing capabilities with respect to inhibiting IL-8 mediated human neutrophil chemotaxis (Figures 50A-50B). The IC<sub>50</sub>'s of 6G4V11N35A IgG1 and 6G4V11N35E IgG1 for human IL-8 were 4.0nM and 6.0nM, respectively, and for rabbit IL-8 were 4.0nM and 2.0nM, respectively. The irrelevant isotype control Fab (4D5) did not inhibit neutrophil migration.

The affinity for IL-8 of these variants relative to the murine 6G4.2.5 mAb was determined using KinExA as described in Section (M). Figure 51 shows the equilibrium constant (Kd) for the full length affinity matured variants 6G4V11N35E IgG1 and 6G4V11N35A IgG1 were approximately 49pM and 88pM, respectively. The Kd for 6G4V11N35A IgG1 was determined directly from the kinetic experiment. As reported with their respective Fabs, this improvement in affinity might be attributed to an approximate 2-fold increase in the on-rate of 6G4V11N35E IgG1 (ka = 3.0x10<sup>6</sup>) compared to that of 6G4V11N35A IgG1 (ka = 8.7x10<sup>5</sup>). In addition, these results were confirmed by a competition radio-immune assay using iodinated human IL-8. 50pM of 6G4V11N35A IgG1 or 6G4V11N35E IgG1 was incubated for 2 hours at 25°C with 30-50pM of <sup>125</sup>I-IL-8 and varying concentrations (0 to 100nM) of unlabeled IL-8. The antibody-antigen mixture was then incubated for 1 hour at 4C with 10ul of a 70% slurry of Protein-A beads (pre-blocked with 0.1% BSA). The beads were briefly spun in a microcentrifuge and the supernatant discarded to remove the unbound <sup>125</sup>I-IL-8. The amount of <sup>125</sup>I-IL-8 specifically bound to the anti-IL-8 antibodies was determined by counting the protein-A pellets in a gamma counter. The approximate Kd values were similar to those determined by KinEXA. The average Kd for 6G4V11N35A IgG1 and 6G4V11N35E IgG1 were 54pM (18 -90pM) and 19pM (5-34pM), respectively (Figure 52).

### 30 T. CONSTRUCTION OF HUMANIZED 6G4V11N35A/E Fab's FOR MODIFICATION BY POLYETHYLENE GLYCOL

A Fab' expression vector for 6G4V11N35A was constructed by digesting p6G4V11N35A.F(ab')<sub>2</sub> with the restriction enzymes Apal and Ndel to remove the 2805 bp fragment encoding the human IgG<sub>1</sub>

constant domain fused with the yeast GCN4 leucine zipper and replacing it with the 2683bp Apal-NdeI fragment from the plasmid pCDNA.18 described in Eigenbrot et al., Proteins: Struct. Funct. Genet., 18: 49-62 (1994). The pCDNA.18 Apal-NdeI fragment carries the coding sequence for the human constant IgG1 heavy domain, including the free cysteine in the hinge region that was used to attach the PEG molecule. The 3758bp Apal-NdeI fragment (encodes the light chain and heavy variable domain of 6G4V11N35A) isolated from p6G4V11N35A.F(ab')<sub>2</sub> was ligated to the 2683bp Apal-NdeI fragment of pCDNA.18 to create p6G4V11N35A.PEG-1. The integrity of the entire coding sequence was confirmed by DNA sequencing. The nucleotide and translated amino acid sequences of heavy chain constant domain with the cysteine in the hinge are presented in Figure 53.

A Fab' expression plasmid for 6G4V11N35E was made similarly by digesting pPH6G4V11N35E (from Section (O) above) with the restriction enzymes Apal and Ndel to isolate the 3758bp Apal-Ndel DNA fragment carrying the intact light chain and heavy variable domain of 6G4V11N35E and ligating it to the 2683 bp Apal-Ndel DNA fragment from p6G4V11N35A.PEG-1 to create p6G4V11N35E.PEG-3. The integrity of the entire coding sequence was confirmed by DNA sequencing.

Anti-IL-8 6G4V11N35A Fab' variant was modified with 20 kD linear methoxy-PEG-maleimide, 30 kD linear methoxy-PEG-maleimide, 40 kD linear methoxy-PEG-maleimide, or 40 kD branched methoxy-PEG-maleimide as described below. All PEG's used were obtained commercially from Shearwater Polymers, Inc.

#### a. MATERIALS AND METHODS

#### Fab'-SH Purification

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A Fab'-SH antibody fragment of the affinity matured antibody 6G4V11N35A was expressed in E. coli grown to high cell density in the fermentor as described by Carter et al., Bio/Technology 10, 163-167 (1992). Preparation of Fab'-SH fragments was accomplished by protecting the Fab'-SH fragments with 4',4'-dithiodipyridine (PDS), partially purifying the protected Fab'-PDS fragments, deprotect the Fab'-PDS with dithiothreitol (DTT) and finally isolate the free Fab'-SH by using gel permeation chromatography.

#### Protection of Fab'-SH with PDS

Fermentation paste samples were dissolved in 3 volumes of 20mM MES, 5mM EDTA, pH 6.0 containing 10.7mg of 4',4'-dithiodipyridine per gram fermentation paste, resulting in a suspension with a pH close to 6.0 The suspension was passed through a homogenizer followed by addition of 5% PEI (w/v), pH 6 to the homogenate to a final concentration of 0.25%. The mixture was then centrifuged to remove solids and the clear supernatant was conditioned to a conductivity of less than 3mS by the addition of cold water.

#### Partial purification of the Fab'-SH molecule using ion exchange chromatography

The conditioned supernatant was loaded onto an ABX (Baker) column equilibrated in 20 mM MES, pH 6.0. The column was washed with the equilibration buffer followed by elution of the Fab'-SH with a 15 column volume linear gradient from 20 mM MES, pH 6.0 to 20 mM MES, 350 mM sodium chloride. The column was monitored by absorbance at 280nm, and the eluate was collected in fractions.

#### Deprotection of the Fab'-SH antibody fragments with DTT

The pH of the ABX pool was adjusted to 4.0 by the addition f dilute HCl. The pH adjusted solution was then deprotected by adding DTT to a final concentration of 0.2mM. The solution was incubated for about 30 minutes and then applied to a gel filtration Sephadex G25 column, equilibrated with 15mM sodium phosphate, 25mM MES, pH 4.0. After elution, the pH of the pool was raised to pH 5.5 and immediately flash frozen at -70°C for storage or derivatized with PEG-MAL as described below.

#### Alternative Fab'-SH Purification

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Alternatively Fab'-SH fragments can be purified using the following procedure. 100 g fermentation paste is thawed in the presence of 200 ml 50 mM acetic acid, pH 2.8, 2 mM EDTA, 1 mM PMSF. After mixing vigorously for 30 min at room temperature, the extract is incubated with 100 mg hen egg white lysozyme. DEAE fast flow resin (approximately 100 mL) is equilibrated with 10 mM MES, pH 5.5, 1 mM EDTA on a sintered glass funnel. The osmotic shock extract containing the Fab'-SH fragment is then filtered through the resin.

A protein G Sepharose column is equilibrated with 10 mM MES, pH 5.5, 1 mM EDTA and then loaded with the DEAE flow-through sample. The column is washed followed by three 4 column volume washes with 10 mM MES, pH 5.5, 1 mM EDTA. The Fab'-SH antibody fragment containing a free thiol is eluted from the column with 100 mM acetic acid, pH 2.8, 1 mM EDTA. After elution, the pH of the pool is raised to pH 5.5 and immediately flash frozen at -70°C for storage or derivatized with PEG-MAL as described below.

#### Preparation of Fab'-S-PEG

The free thiol content of the Fab'-SH preparation obtained as described above was determined by reaction with 5,5'-dithiobis(2-nitrobenzoic acid) (DTNB) analysis according to the method of Creighton in Protein Structure: A Practical Approach, Creighton, T.E., ed, IRL Press (Oxford, UK: 1990), pp. 155-167. The concentration of free thiol was calculated from the increase on absorbance at 412 nm, using  $e_{412} = 14,150 \text{ cm}^{-1} \text{ M}^{-1}$  for the thionitrobenzoate anion and a  $M_r = 48,690$  and  $e_{280} = 1.5$  for the Fab'-SH antibody. To the Fab'-SH protein G Sepharose pool, or the deprotected Fab'-SH gel permeation pool, 5 molar equivalents of PEG-MAL were added and the pH was immediately adjusted to pH 6.5 with 10% NaOH.

The Fab'-S-PEG was purified using a 2.5 x 20 cm cation exchange column (Poros 50-HS). The column was equilibrated with a buffer containing 20 mM MES, pH 5.5. The coupling reaction containing the PEGylated antibody fragment was diluted with deionized water to a conductivity of approximately 2.0 mS. The conditioned coupling reaction was then loaded onto the equilibrated Poros 50 HS column. Unreacted PEG-MAL was washed from the column with 2 column volumes of 20 mM MES, pH 5.5. The Fab'-S-PEG was eluted from the column using a linear gradient from 0 to 400 mM NaCl, in 20 mM MES pH 5.5, over 15 column volumes.

Alternatively a Bakerbond ABX column can be used to purify the Fab'-S-PEG molecule. The column is equilibrated with 20 mM MES, pH 6.0 (Buffer A). The coupling reaction is diluted with deionized water until the conductivity equaled that of the Buffer A (approximately 2.0 mS) and loaded onto the column. Unreacted PEG-MAL is washed from the column with 2 column volumes of 20 mM MES, pH 6.0. The Fab'-S-PEG is eluted from the column using a linear gradient from 0 to 100 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, in 20 mM MES pH 6.0, over 15 column volumes.

#### Size Exclusion Chromatography

The hydrodynamic or effective size of each molecule was determined using a Pharmacia Superose-6 HR 10/30 column (10x300mm). The mobile phase was 200 mM NaCl, 50 mM sodium phosphate pH 6.0. Flow rate was at 0.5 ml/min and the column was kept at ambient temperature. Absorbance at 280 nm was monitored where PEG contributed little signal. Biorad MW standards containing cyanocobalamin, myoglobin, ovalbumin, 1gG, Thyroglobulin monomer and dimer were used to generate a standard curve from which the effective size of the pegylated species was estimated.

#### b. RESULTS

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#### Size Exclusion Chromatography

The effective size of each modified species was characterized using size exclusion chromatography. The results are shown in Fig. 60 below. The theoretical molecular weight of the anti-IL8 Fab fragments modified with PEG 5kD, 10kD, 20kD, 30kD, 40kD (linear), 40kD (branched) or 100,000kD is shown along with the apparent molecular weight of the PEGylated fragments obtained by HPLC size exclusion chromatography. When compared to the theoretical molecular weight of the Fab'-S-PEG fragments, the apparent molecular weight (calculated by size exclusion HPLC) increases dramatically by increasing the size of the PEG attached to the fragments. Attachment of a small molecular weight PEG, for example PEG 10,000D only increases the theoretical molecular weight of the PEGylated antibody fragment (59,700 D) by 3 fold to an apparent molecular weight of 180,000D. In contrast attachment of a larger molecular weight PEG for example 100,000D PEG to the antibody fragment increases the theoretical molecular weight of the PEGylated antibody fragment (158,700 D) by 12 fold to an apparent molecular weight of 2,000,000D.

#### SDS-PAGE

In Fig. 61, the upper panel shows the size of the anti-IL-8 Fab fragments modified with PEG of molecular weight 5kD (linear), 10kD (linear), 20kD (linear), 30kD (linear), 40kD (linear), 40kD (branched) or 100kD (linear) under reduced conditions. The unmodified Fab is shown in lane 2 from right to left. Both the heavy and light chains of the Fab had a molecular weight of approximately 30kD as determined by PAGE. Each PEGylated fragment sample produced two bands: (1) a first band (attributed to the light chain) exhibiting a molecular weight of 30kD; and (2) a second band (attributed to the heavy chain to which the PEG is attached specifically at the hinge SH) exhibiting increasing molecular weights of 40, 45, 70, 110, 125, 150 and 300kD. This result suggested that PEGylation was specifically restricted to the heavy chain of the Fab's whereas the light chain remained unmodified.

The lower panel is non-reduced PAGE showing the size of the anti-IL-8 Fab fragments modified with PEG of molecular weight 5kD (linear), 20kD (linear), 30kD (linear), 40kD (linear), 40kD (branched), or 100kD (linear). The PEGylated fragments exhibited molecular weights of approximately 70kD, 115kD, 120kD, 140kD, 200kD and 300kD.

The SDS PAGE gels confirm that all Fab'-S-PEG molecules were purified to homogeneity and that the molecules differed only with respect to the size of the PEG molecule attached to them.

#### U. AMINE SPECIFIC PEGYLATION OF ANTI-IL-8 F(ab')<sub>2</sub> FRAGMENTS

Pegylated F(ab')<sub>2</sub> species were generated by using large MW or branched PEGs in order to achieve a large effective size with minimal protein modification which might affect activity. Modification involved N-hydroxysuccinamide chemistry which reacts with primary amines (lysines and the N-terminus). To decrease the probability of modifying the N-terminus, which is in close proximity to the CDR region, a reaction pH of 8, rather than the commonly used pH of 7, was employed. At pH 8.0, the amount of the reactive species (charged NH<sub>3</sub>\*) would be considerably more for the ε-NH2 group of lysines (pK<sub>a</sub>=10.3) than for the α-NH2 group (pK<sub>a</sub> of approximately 7) of the amino-terminus. For the linear PEGs, a methoxy-succinimidyl derivative of an NHS-PEG was used because of the significantly longer half-life in solution (17 minutes at 25°C at pH 8.0) compared to the NHS esters of PEGs (which have 5-7 minute half life under the above conditions). By using a PEG that is less prone to hydrolysis, a greater extent of modification is achieved with less PEG. Branched PEGs were used to induce a large increase in effective size of the antibody fragments.

#### a. MATERIALS

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All PEG reagents were purchased from Shearwater Polymers and stored at -70°C in a desiccator: branched N-hydroxysuccinamide-PEG (PEG2-NHS-40KDa) has a 20 kDa PEG on each of the two branches, methoxy-succinimidyl-propionic acid-PEG (M-SPA-20000) is a linear PEG molecule with 20 kDa PEG. Protein was recombinantly produced in *E. coli* and purified as a (Fab)'<sub>2</sub> as described in Sections (K) and (O) above.

#### b. METHODS

IEX method: A J. T. Baker Wide-Pore Carboxy-sulfone (CSX), 5 micron, 7.75 x 100 mm HPLC column was used for fractionation of the different pegylated products, taking advantage of the difference in charge as the lysines are modified. The column was heated at 40°C. A gradient as shown in Table 7 below was used where Buffer A was 25 mM sodium Borate/25 mM sodium phosphate pH 6.0, and Buffer B was 1 M ammonium sulfate, and Buffer C was 50 mM sodium acetate pH 5.0.

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Table 7

5	Time (min)	%B	%С	flow mL/min
	0	10	10	1.5
	20	18	7.5	1.5
	25	25	7.5	1.5
10	27	70	3.0	2.5
• •	29	70	3.0	2.5
	30	10	10	2.5
	33	10	10	2.5

SEC-HPLC: The hydrodynamic or effective size of each molecule was determined using a Pharmacia Superose-6 HR 10/30 column (10x300mm). The mobile phase was 200 mM NaCl, 50 mM sodium phosphate pH 6.0. Flow rate was at 0.5 ml/min and the column was kept at ambient temperature. Absorbance at 280 nm was monitored where PEG contributed little signal. Biorad MW standards containing cyanocobalamin, myoglobin, ovalbumin, IgG, Thyroglobulin monomer and dimer were used to generate a standard curve from which the effective size of the pegylated species was estimated.

SEC-HPLC-Light Scattering: For determination of the exact molecular weight, this column was connected to an on-line light scattering detector (Wyatt Minidawn) equipped with three detection angles of 50°, 90°, and 135° C. A refractive index detector (Wyatt) was also placed on-line to determine concentration. All buffers were filtered with Millipore 0.1  $\mu$  filters; in addition al 0.02  $\mu$  Whatman Anodisc 47 was placed on-line prior to the column.

The intensity of scattered light is directly proportional to the molecular weight (M) of the scattering species, independent of shape, according to:

$$M = R_0/K \cdot c$$

where  $R_0$  is the Rayleigh ratio, K is an optical constant relating to the refractive index of the solvent, the wavelength of the incident light, and dn/dc, the differential refractive index between the solvent and the solute with respect to the change in solute concentration, c. The system was calibrated with toluene ( $R_0$  of  $1.406 \times 10^{-5}$  at 632.8 nm); a dn/dc of 0.18, and an extinction coefficient of 1.2 was used. The system had a mass accuracy of ~5%.

SDS-PAGE: 4-12% Tris-Glycine Novex minigels were used along with the Novex supplied Tris-Glycine running buffers. 10-20 ug of protein was applied in each well and the gels were run in a cold box at 150 mV/gel for 45 minutes. Gels were then stained with colloidal Coomassie Blue (Novex) and then washed with water for a few hours and then preserved and dried in drying buffer (Novex)

Preparati n of a linear(1)20KDa-(N)-(Fab')2: A 4 mg/ml solution of anti-IL8 formulated initially in a pH 5.5 buffer was dialyzed overnight against a pH 8.0 sodium phosphate buffer. 5 mL protein

was mixed at a molar ratio of 3:1. The reaction was carried out in a 15mL polypropylene Falcon tube and the PEG was added while vort xing the sample at low speed for 5 seconds. It was then placed on a nutator for 30 minutes. The extent of modification was evaluated by SDS-PAGE. The whole 5 ml reaction mixture was injected on the IEX for removal of any unreacted PEG and purification of singly or doubly pegylated species. The above reaction generated a mixture of 50% singly-labeled anti-IL8. The other 50% unreacted anti-IL8 was recycled through the pegylation/purification steps. The pooled pegylated product was dialyzed against a pH 5.5 buffer for in vitro assays and animal PK studies. Endotoxin levels were measured before administration to animals or for the cell based assays. Levels were below 0.5 eu/ml. The fractions were also run on SDS-PAGE to confirm homogeneity. Concentration of the final product was assessed by absorbance at 280 nm using an extinction coefficient of 1.34, as well as by amino acid analysis.

Preparation of a branched(1)40KDa-(N)-(Fab')2: A 4 mg/mL solution of anti-IL8 (Fab')2 formulated in a pH 5.5 buffer was dialyzed overnight against a pH 8.0 phosphate buffer. Solid PEG powder was added to 5 mL protein in two aliquots to give a final PEG:protein molar ratio of 6:1. Each solid PEG aliquot was added to the protein in a 15 mL polypropylene Falcon tube while vortexing at low speed for 5 sec, and then placing the sample on a nutator for 15 minutes. The extent of modification was evaluated by SDS-PAGE using a 4-12% Tris-Glycine (Novex) gel and stained with colloidal Coomasie blue (Novex). The 5 mL PEG-protein mixture was injected on the ion exchange column for removal of any unreacted PEG. The above reaction generated a mixture of unreacted (37%), singly-labelled (45%), doubly and triply-labeled (18%) species. These were the optimal conditions for obtaining the greatest recovery of the protein with only 1 PEG per antibody rather than the higher molecular weight adducts. The unmodified anti-IL8 was recycled. The pegylated products were separated and fractionated in falcon tubes and then dialyzed against a pH 5.5 buffer for assays and animal PK studies. Endotoxin levels were below 0.5 eu/ml. The fractions were also run on SDS-PAGE to confirm homogeneity. The concentration of the final product was assessed by absorbance at 280 nm using an extinction coefficient of 1.34, as well as by amino acid analysis.

Preparation of branched(2)-40KDa-(N)(Fab')2: This molecule was most efficiently made by adding three times in 15 minute intervals a 3:1 molar ratio of PEG to the already modified branched(1)-40KDa-(N)-(Fab')2. The molecule was purified on IEX as 50% branched(2)-40KDa-(N)-(Fab')2. The unmodified molecule was recycled until ~20 mg protein was isolated for animal PK studies. The product was characterized by SEC-light scattering and SDS-PAGE.

#### c. RESULTS

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PEGs increased the hydrodynamic or effective size of the product significantly as determined by gel filtration (SEC-HPLC). Figure 62 shows the SEC profile of the pegylated F(ab')<sub>2</sub> species with UV detection at 280 nm. The hydrodynamic size of each molecule was estimated by reference to the standard MW calibrators. As summarized in Figure 62, the increase in the effective size of (Fab')<sub>2</sub> was about 7-fold

by adding one linear 20 kDa PEG molecule and about 11-fold by adding one branched ("Br(1)") 40 kDa PEG molecule, and somewhat more with addition of two branched ("Br(2)") PEG m lecules.

Light scattering detection gave the exact molecular weight of the products and confirmed the extent of modification (Figure 63). The homogeneity of the purified material was shown by SDS-PAGE (Figure 64). Underivatized F(ab')<sub>2</sub> migrated as a 120 kDa species, the linear(1)20KD-(N)-F(ab')<sub>2</sub> migrated as a band at 220kDa, the Br(1)-40KD(N)-F(ab')<sub>2</sub> migrated as one major band at 400 kDa, and the Br(2)-40KD-(N)-F(ab')<sub>2</sub> migrated as a major band at around 500 kDa. The proteins appeared somewhat larger than their absolute MW due to the steric effect of PEG.

### V. <u>IN VITRO ACTIVITY CHARACTERIZATION OF PEG MODIFIED Fab' FRAGMENTS OF 6G4V11N35A (MALEIMIDE CHEMICAL COUPLING METHOD)</u>

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Anti-IL-8 6G4V11N35A Fab' variants modified with 5-40kD linear PEG molecules and a 40kD branched PEG molecule were tested for their ability to inhibit both IL-8 binding and activation of human neutrophils; the procedures were described in Sections (B)(1), (B)(2) and (B)(3) above. The binding curves and IC<sub>50</sub>'s for PEG-maleimide modified 6G4V11N35A Fab' molecules are presented in Figures 54A-54C. The IC<sub>50</sub> of the 5kD pegylated Fab' (350pM) and the average IC<sub>50</sub> of the Fab control (366pM) were not significantly different, suggesting that the addition of a 5kD MW PEG did not affect the binding of IL-8 to the modified Fab' (Figure 54A). However, a decrease in the binding of IL-8 to the 10kD and 20kD pegylated Fab' molecules was observed as depicted by the progressively higher IC<sub>50</sub>'s (537pM and 732pM, respectively) compared to the average IC<sub>50</sub> of the native Fab. These values represent only a minimal loss of binding activity (between 1.5- and 2.0-fold). A less pronounced difference in IL-8 binding was observed for the 30kD and 40kD linear PEG antibodies (Figure 54B). The IC<sub>50</sub>'s were 624pM and 1.1nM, respectively, compared to the 802pM value of the Fab control. The 40kD branched PEG Fab' showed the largest decrease in IL-8 binding (2.5 fold) relative to the native Fab (Figure 54C). Nevertheless, the reduction in binding of IL-8 by these pegylated Fab's is minimal.

The ability of the pegylated antibodies to block IL-8 mediated activation of human neutrophils was demonstrated using the PMN chemotaxis (according to the method described in Section B(2) above) and β-glucuronidase release (according to the method described in Lowman et al., J. Biol. Chem., 271: 14344 (1996)) assays. The IC<sub>50</sub>'s for blocking IL-8 mediated chemotaxis are shown in Figures 55A-55C. The 5-20kD linear pegylated Fab' antibodies were able to block IL-8 mediated chemotaxis within 2-3 fold of the unpegylated Fab control (Figure 55A). This difference is not significant because the inherent variation can be up to 2 fold for this type of assay. However, a significant difference was detected for the 30kD and 40kD linear pegylated Fab' antibodies as illustrated by the higher IC<sub>50</sub>'s of the 30kD linear PEG-Fab' (2.5nM) and 40kD linear PEG-Fab' (3.7nM) compared to the Fab control (0.8nM) (Figure 55B).

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The ability of the 40kD branched PEG Fab' molecule to block IL-8 mediated chemotaxis was similar to that of the 40kD linear PEG Fab' (Figure 55C). At most, the ability of the pegylated Fab' antibodies to block IL-8 mediated chemotaxis was only reduced 2-3 fold. Furthermore, release of β-glucuronidase from the granules of neutrophils was used as another criteria for assessing IL-8 mediated activation of human PMNs. Figure 56A (depicting results obtained with 5 kD, 10 kD and 20 kD linear PEGs), Figure 56B (depicting results obtained with 30 kD and 40 kD linear PEGs), and Figure 56C (depicting results obtained with 40 kD branched PEG) show that all the pegylated Fab' antibodies were able to inhibit IL-8 mediated release of β-glucuronidase as well as or better than the unpegylated Fab control. The data collectively shows that the pegylated Fab' variants are biological active and are capable of inhibiting high amounts of exogenous IL-8 in in-vitro assays using human neutrophils.

### W. <u>IN VITRO ACTIVITY CHARACTERIZATION OF PEG MODIFIED F(ab')</u> FRAGMENTS OF 6G4V1 IN35A (SUCCINIMIDYL CHEMICAL COUPLING METHOD)

The anti-IL-8 variant 6G4V11N35A F(ab')<sub>2</sub> modified with (a) a single 20kD linear PEG molecule per F(ab')<sub>2</sub>, (b) a single 40kD branched PEG molecule per F(ab')<sub>2</sub>, (c) with three, four, or five 20 kD linear PEG molecules per F(ab')<sub>2</sub>; (a mixture of: (1) species having three 20 kD linear PEG molecules per F(ab')<sub>2</sub>; (2) species having four 20 kD linear PEG molecules per F(ab')<sub>2</sub>; and (3) species having five 20 kD linear PEG molecules per F(ab')<sub>2</sub>; denoted as "20 kD linear PEG (3,4,5) F(ab')<sub>2</sub>"), or (d) with two 40kD branched PEG molecules per F(ab')<sub>2</sub> (denoted as "40 kD branch PEG (2) F(ab')<sub>2</sub>"), were tested for their ability to inhibit <sup>125</sup>1-IL-8 binding and to neutralize activation of human neutrophils. The procedures used are described in Sections (B)(1), (B)(2) and (B)(3) above. The binding curves for pegylated F(ab')<sub>2</sub> variants are shown in Figures 57A-57B. No significant differences were observed amongst the F(ab')<sub>2</sub> control, the single 20kD linear PEG-modified F(ab')<sub>2</sub>, and the single 40kD branched PEG-modified F(ab')<sub>2</sub> (Figure 57A). However, the F(ab')<sub>2</sub> variants containing multiple PEG molecules showed a slight reduction (less than 2-fold) in their ability to bind IL-8. The IC<sub>50</sub>'s of the 20kD linear PEG (3,4,5) F(ab')<sub>2</sub> and 40kD branch PEG (2) F(ab')<sub>2</sub> variants were 437pM and 510pM, respectively, compared to 349pM of the F(ab')<sub>2</sub> control (Figure 57B).

The ability of these pegylated F(ab')<sub>2</sub> variants to block IL-8 mediated neutrophil chemotaxis is presented in Figures 58A-58B. Consistent with the PMN binding data, the single linear and branched PEG F(ab')<sub>2</sub> variants were able to block IL-8 mediated chemotaxis similar to the unpegylated F(ab')<sub>2</sub> control (Figure 58A). The ability of the 40kD branch PEG (2) F(ab')<sub>2</sub> variant to inhibit PMN chemotaxis was

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identical to the control F(ab')<sub>2</sub> while the 20kD linear PEG (3,4,5) F(ab')<sub>2</sub> mixture was able to inhibit within 3-fold of the control antibody (Figure 58B).

Shown in Figures 59A and 59B are the results of the  $\beta$ -glucuronidase release assay which is a measure of degranulation by IL-8 stimulated human neutrophils. The single 20kD linear PEG-modified  $F(ab')_2$  and the single 40kD branched PEG-modified  $F(ab')_2$  variants were able to inhibit release of  $\beta$ -glucuronidase as well as the  $F(ab')_2$  control (Figure 59A). The 40kD branch PEG (2)  $F(ab')_2$  inhibited this response within 2-fold of the  $F(ab')_2$  control (Figure 59B). The 20kD linear PEG (3,4,5) molecule was not tested. Overall, the  $F(ab')_2$  pegylated anti-IL-8 antibodies were biologically active and effectively prevented IL-8 binding to human neutrophils and the signaling events leading to cellular activation.

# 10 X. PHARMACOKINETIC AND SAFETY STUDY OF EIGHT CONSTRUCTS OF PEGYLATED ANTI-IL-8 (HUMANIZED) F(AB')2 AND FAB' FRAGMENTS IN NORMAL RABBITS FOLLOWING INTRAVENOUS ADMINISTRATION

The objective of this study was to evaluate the effect of pegylation on the pharmacokinetics and safety of six pegylated humanized anti-IL-8 constructs (pegylated 6G4V11N35A.Fab' and pegylated 6G4V11N35A.F(ab')<sub>2</sub> obtained as described in Sections (T) and (U) above) relative to the non-pegylated fragments in normal rabbits. Eight groups of two/three male rabbits received equivalent protein amounts of pegylated 6G4V11N35A.Fab' or pegylated 6G4V11N35A.F(ab')<sub>2</sub> constructs (2 mg/kg) via a single intravenous (IV) bolus dose of one anti-IL8 construct. Serum samples were collected according to the schedule shown in Table 8 below and analyzed for anti-IL8 protein concentrations and antibody formation against anti-IL8 constructs by ELISA.

Table 8

Group No.	Dose level/ Route	Material	Blood Collection
1		Fab' control	0,5,30 min; 1,2,3,4,6,8,10, 14,20,24,360 hr
2	!	linear(1)20K(s)Fab'	
3		linear(1)40K(s)Fab'	0,5,30 min; 1,2,4,6,8,10,12, 24,28,32,48,72,96,168,216,
4	2 mg/kg	branched(1)40K(N)F(ab') <sub>2</sub>	264,336,360 hr
5	(protein conc.) IV bolus	F(ab') <sub>2</sub> control	0,5,30 min; 1,2,4,6,8,10,12, 24,28,32,48,52,56,336 hr

Group No.	Dose level/ Route	Material	Blood Collection
6		branched(2)40K(s)Fab'	0,5,30 min; 1,2,4,6,8,10,12, 24,28,32,48,72,96,168,216,264,3 36 hr; Day 17,21, 25
7		branched(2)40K(N)F(ab') <sub>2</sub>	0,5,30 min; 1,2,4,6,8,10,12, 24,28,32,48,72,144,192, 240 hr; Day 13, 16, 20, 23
8		linear(1)30K(s)Fab'	0,5,30 min; 1,2,4,6,8,10,12, 24,28,32,48,72,96,168,216,264,3 36 hr; Day 17,21, 25

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#### a. METHODS

Three male New Zealand White (NZW) rabbits per group (with exception to Group 7, n=2) received an equivalent amount of 6G4V11N35A protein (Fab' or F(ab')<sub>2</sub>) construct at 2 mg/kg via an IV bolus dose in a marginal ear vein. Amino acid composition analysis and absorbance at 280 nm using extinction coefficients of 1.26 for 6G4V11N35A Fab' constructs and 1.34 for 6G4V11N35A F(ab')<sub>2</sub> constructs were performed to determine the protein concentration. Whole blood samples were collected via an ear artery cannulation (ear opposing dosing ear) at the above time points. Samples were harvested for serum and assayed for free 6G4V11N35A Fab' or F(ab')<sub>2</sub> constructs using an IL-8 Binding ELISA. Assays were conducted throughout the study as samples became available. All animals were sacrificed following the last blood draw, and necropsies were performed on all animals in Groups 1, 4–8. Due to the development of antibodies against the 6G4V11N35A constructs, non-compartmental pharmacokinetic analysis was conducted on concentration versus time data only up to 168 hours.

#### b. RESULTS

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In four animals (Animals B, P, Q, V), interference to rabbit serum in the ELISA assay was detected (i.e. measurable concentrations of anti-IL8 antibodies at pre-dose). However, because these values were at insignificant levels and did not effect the pharmacokinetic analysis, the data were not corrected for this interference.

One animal (Animal G; Group 3) was exsanguinated before the termination of the study and was excluded from the pharmacokinetic analysis. At 4 hours, the animal showed signs of a stroke that was not believed to be drug related, as this can occur in rabbits following blood draws via ear artery cannulation.

The mean concentration-time profiles of the eight anti-IL8 constructs in normal rabbits are depicted in Fig. 65, and the pharmacokinetic parameters for the eight constructs are summarized in Table 9 below. Significant antibodies to the anti-IL-8 constructs were present at Day 13/14 in all dose groups except Group 1 (Fab' control).

Table 9. Pharmacokinetic parameters.

Molecule	Fab'					F(ab') <sub>2</sub>		
Group No.	1	2	8	3	6	5	4	7
PEG structure		linear	linear	linear	branched	_	branched	branched
Number of PEGs	_	1	ì	1	1	-	1	2
PEG MW		20K	30K	40K	40K		40K	40K
Dose (mg/kg)	2	2	2	2	2	2	2	2
V <sub>c</sub> (mL/kg)	58±3	36±3	35±1	34	44±1	45±5	36±1	32
V <sub>ss</sub> (mL/kg)	68±8	80±8	110±15	79	88±21	59±4	50±3	52
Cmax (µg/mL)	35±1	58±3	57±1	60	45±1	45±6	56±2	62
Tmax (min)	5	5	5	5	5	5	5	5
t <sub>1/2</sub> term (hr)	3.0±0.9	44±2	43±7	50	105±11	8.5±2.1	45±3	48
AUC <sub>0-</sub> (hr•µg/mL)	18±3	80±74	910±140	1600	3400±1300	140±3	2200±77	2500
CL (mL/hr/kg) g	110±17	2.5±0.2	2.2±0.4	1.3	0.63±0.20	14±0	0.92±0.03	0.83
MRT (hr) h	0.61±0.15	32±2	45±9	63	140±18	4.2±0.3	55±3	64
No. of Animals	3	3_	3	2	3	3	3	2

Initial volume of distribution.

The initial volume of distribution approximated the plasma volume for both the Fab' and F(ab')2.

Pegylation decreased serum CL of anti-IL8 fragments and extended both the terminal half-life and MRT as shown in Table 10 below.

Table 10. Fold decrease/increase in clearance, terminal half-life & MRT of pegylated anti-IL8 fragments.

anti-IL8 fragment Group No.		Fab'	Fab'				F(ab'	F(ab') <sub>2</sub>		
		1	2	8	3	6	5	4	7	
PEG struct No. of PEC	ure	-	linear I 20K	linear 1 30K	linear 1 40K	bran. 1 40K	-	bran. 1 40K	bran. 2 40K	
PEG MW		1110	2.5	2.2	1.3	0.63	14	0.92	0.83	
CL:	mean (mL/hr/kg) fold decrease	110	46	51	90	180	1	15	17	
t1/2 term :	mean (hr) fold increase	3.0	44 14	43 14	50 17	110 35	8.5 1	45 5.3	48 5.7	
MRT:	mean (hr) fold increase	0.61	32 53	45 73	63 100	140 240	4.2	55 13	64 15	

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Volume of distribution at steady state.

Observed maximum concentration.

Observed time to Cmax.

t<sub>1/2</sub> term= half-life associated with the terminal phase of the concentration vs. time profile.

Area under the concentration versus time curve (extrapolated to infinity).

CL= serum clearance.

MRT= Mean residence time.

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For the pegylated anti-IL8 Fab' fragments, CL decreased by 46 to 180-fold. Terminal half-life and MRT increased 14 to 35-fold and 53 to 240-fold, respectively. For pegylated anti-IL8 F(ab')<sub>2</sub> molecules, CL decreased 15 to 17-fold with pegylation, and terminal half-life and MRT increased by greater than 5-fold and 13-fold, respectively. The changes in these parameters increased for both pegylated Fab' and F(ab')<sub>2</sub> molecules with increasing PEG molecular weight and approached the values of the full-length anti-IL8 (terminal half-life of 74 hours, MRT of 99 hours and CL of 0.47 mL/hr/kg). In comparing the branched(1)40K Fab' (Group 6) and branched(1)40K F(ab')<sub>2</sub> (Group 4), unexpected pharmacokinetics were observed. The pegylated Fab' molecule appeared to remain in the serum longer than the pegylated F(ab')<sub>2</sub> (see Figure 66). The mean CL of branched(1)40K Fab' was 0.63 mL/hr/kg, but a higher CL was observed for branched(1)40kD F(ab')<sub>2</sub> (CL 0.92 mL/hr/kg). The terminal half-life, likewise, was longer for the Fab' than the F(ab')<sub>2</sub> pegylated molecule (110 vs 45 hours).

The pharmacokinetic data demonstrated that pegylation decreased CL and increased terminal t1/2 and MRT of anti-IL8 fragments (Fab' and F(ab')<sub>2</sub>) to approach that of the full-length anti-IL8. Clearance was decreased with pegylation 46 to 180-fold for the Fab' and approximately 16-fold for the F(ab')<sub>2</sub>. The terminal half-life of the Fab' anti-IL8 fragment was increased by 14 to 35-fold and approximately 5-fold for the F(ab')<sub>2</sub> anti-IL8. MRT, likewise, were extended by 53 to 240-fold for the Fab' and approximately 14-fold for the F(ab')<sub>2</sub>. The branched(1) 40kD Fab' had a longer terminal half-life and lower clearance compared to the branched(1) 40kD F(ab')<sub>2</sub>.

## Y. <u>IN VIVO EFFICACY TESTING OF ANTI-IL-8 ANTIBODY REAGENTS IN RABBIT MODEL</u> OF ISCHEMIA/REPERFUSION AND ACID ASPIRATION-INDUCED ACUTE RESPIRATORY DISTRESS SYNDROME (ARDS)

Full length murine anti-rabbit IL-8 monoclonal antibody 6G4.2.5, 40 kD branched PEG-6G4V11N35A Fab', and control antibody (anti-HIV gp120 monoclonal antibody 9E3.1F10) were tested in a rabbit ARDS model. The animals were weighed and anaesthetized by intramuscular injection of ketamine (50 mg/kg body weight), xylazine (5 mg/kg body weight), and acepromazine (0.75 mg/kg body weight). A second dose (20% of the first dosage) was given IM 15 minutes before removal of vascular clip, and third dose (60% of the first dosage) was given at tracheotomy. Intra-arterial catheter (22G, 1 in. Angiocath) and intra-venous catheter (24G, 1 in. angiocath) were be placed in the ear central artery and posterior marginal ear vein for blood samplings (arterial blood gases and CBC) and anti-IL-8 and fluid administration, respectively. The anaesthetized animals were transferred in a supine position to an operating tray; the abdominal area was shaved and prepared for surgery. Via a midline laparotomy, the superior mesenteric artery (SMA) was isolated and a microvascular arterial clip applied at the aortic origin. Before the temporary closure of the abdomen using 9 mm wound clip (Autoclip, Baxter), 15 ml of normal saline was

given intraperitoneally as fluid supplement. After 110 minutes of intestinal ischemia, the abdominal incision was reopened and the arterial clip was released to allow reperfusion. Before closure, 5 ml of normal saline was given intraperitoneally for fluid replacement. The laparotomy incision was closed in two layers and the animals allowed to awaken.

After surgery, the animals were placed on a heating pad (38°C) and continuously monitored for up to 6 hours post reperfusion and lactated Ringer's 8-12 ml/kg/hr IV was given as fluid supplement.

At 22-24 hr post-reperfusion, a tracheotomy was performed under anesthesia. Normal physiologic saline was diluted 1:3 with water and adjusted to pH 1.5 (adjusted by using 1N HCL); 3 ml/kg body weight was then instilled intra-tracheally. Rectal temperature was maintained at 37 +/- 1 degree C using a homeothermic heat therapy pad (K-Mod II, Baxter). Fluid supplements (LRS) at a rate of 5 ml/kg/hour IV were given. Blood gases were monitored every hour. The rabbits were returned to the cage after 6 hr of continuous monitoring.

Just prior to aspiration, animals were treated with saline, the control monoclonal antibody (anti-HIV gp-120 IgG 9E3.1F10), the full length murine anti-rabbit IL8 (6g4.2.5 murine IgG2a anti-rabbit IL8) or the pegylated 6G4V11N35A Fab' (6G4V1N35A Fab' modified with 40kD branched PEG-maleimide as described in Section T above, denoted as "40 kD branched PEG-6G4V11N35A Fab' "). Data from saline or control antibody treated animals was combined and presented as "Control". Arterial blood gases and A-a PO2 gradient measurements were taken daily, and IV fluid supplementation was performed daily. A-a PO2 gradient was measured at 96 hr of reperfusion. The A-a PO2 gradient was calculated as:

A-a PO2 = [FIO2(PB - PH2O) - (PaCO2/RQ)] - PaO2.

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PaO2/FiO2 ratios were measured at 24hr and 48hr in room air and 100% oxygen.

After the final A-a PO2 gradient measurement, the animals were anesthetized with Nembutal 100mg/kg i.v. and the animals were euthanized by transecting the abdominal aorta in order to reduce red blood cell contamination of bronchoalveolar lavage fluid (BAL). The lungs were removed en bloc. The entire lung was weighed and then lavaged with an intratracheal tube (Hi-Lo tracheal tube, 3mm) using 30 ml of HBSS and lidocain. Total and differential leukocyte counts in the BAL were determined. Lesions/changes were verified by histological examination of each lobe of the right lung of each animal.

The gross lung weight, total leukocyte and polymorphonuclear cell counts in BAL, and PaO2/FiO2 data obtained are depicted in Figs. 67, 68 and 69, respectively. Treatment with 40 kD branched PEG-6G4V11N35A Fab' exhibited no effect on the biological parameters measured in the model as compared to the "Control" group. However, the data do not contradict the pharmacokinetic analysis or the in vitro activity analysis for the 40 kD branched PEG-6G4V11N35A Fab' presented in Sections (V) and (X) above. In addition, these data do not contradict the ability of the 40 kD branched PEG-6G4V11N35A Fab' to reach and act on disease effector targets in circulation or other tissues.

The following biological materials have been deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD, USA (ATCC):

	Material	ATCC Accession No.	Deposit Date
	hybridoma cell line 5.12.14	HB 11553	February 15, 1993
	hybridoma cell line 6G4.2.5	HB 11722	September 28, 1994
5	pantiIL-8.2, E. coli strain 294 mm	97056	February 10, 1995
	p6G425chim2, E. coli strain 294 mm	97055	February 10, 1995
	p6G4V11N35A.F(ab') <sub>2</sub>	97890 .	February 20, 1997
	E. coli strain 49D6(p6G4V11N35A.F(ab') <sub>2</sub>	98332	February 20, 1997
	p6G425V11N35A.choSD	209552	December 16, 1997
10	clone#1933 alL8.92 NB 28605/12	CRL-12444	December 11, 1997
	clone#1934 aIL8.42 NB 28605/14	CRL-12445	December 11, 1997

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These deposits were made under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purpose of Patent Procedure and the Regulations thereunder (Budapest Treaty). This assures maintenance of a viable deposit for 30 years from the date of deposit. These cell lines will be made available by ATCC under the terms of the Budapest Treaty, and subject to an agreement between Genentech, Inc. and ATCC, which assures permanent and unrestricted availability of the cell lines to the public upon issuance of the pertinent U.S. patent or upon laying open to the public of any U.S. or foreign patent application, whichever comes first, and assures availability of the cell lines to one determined by the U.S. Commissioner of Patents and Trademarks to be entitled thereto according to 35 USC §122 and the Commissioner's rules pursuant thereto (including 37 CFR §1.14 with particular reference to 886 OG 638).

The assignee of the present application has agreed that if the deposited cell lines should be lost or destroyed when cultivated under suitable conditions, they will be promptly replaced on notification with a specimen of the same cell line. Availability of the deposited cell lines is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent laws

#### SEQUENCE LISTING

(1) GENERAL INFORMATION: 5 (i) APPLICANT: Hsei, Vanessa Koumenis, Iphigenia Leong, Steven R. Presta, Leonard G. Shahrokh, Zahra 10 Zapata, Gerardo A. (ii) TITLE OF INVENTION: Antibody Fragment-Polymer Conjugates and Humanized Anti-IL-8 Monoclonal Antibodies 15 (iii) NUMBER OF SEQUENCES: 76 (iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: Genentech, Inc. (B) STREET: 1 DNA Way 20 (C) CITY: South San Francisco (D) STATE: California (E) COUNTRY: USA (F) ZIP: 94080 25 (v) COMPUTER READABLE FORM: (A) MEDIUM TYPE: 3.5 inch, 1.44 Mb floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: WinPatin (Genentech) 30 (vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: 20-Feb-1998 (C) CLASSIFICATION: 35 (viii) ATTORNEY/AGENT INFORMATION: (A) NAME: Love, Richard B. (B) REGISTRATION NUMBER: 34,659 (C) REFERENCE/DOCKET NUMBER: P1085R3PCT 40 (ix) TELECOMMUNICATION INFORMATION: (A) TELEPHONE: 650/225-5530 (B) TELEFAX: 650/952-9881 (2) INFORMATION FOR SEQ ID NO:1: 45 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: Nucleic Acid 50 (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

## CAGTCCAACT GTTCAGGACG CC 22

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- (2) INFORMATION FOR SEQ ID NO:2:
- 5 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 22 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
  - (D) TOPOLOGY: Linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

# GTGCTGCTCA TGCTGTAGGT GC 22

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- (2) INFORMATION FOR SEQ ID NO:3:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 23 base pairs
- (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
  - (D) TOPOLOGY: Linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

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## GAAGTTGATG TCTTGTGAGT GGC 23

(2) INFORMATION FOR SEQ ID NO:4:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 24 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

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(D) TOPOLOGY: Linear

- 40 GCATCCTAGA GTCACCGAGG AGCC 24
  - (2) INFORMATION FOR SEQ ID NO:5:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22 base pairs
      - (B) TYPE: Nucleic Acid
      - (C) STRANDEDNESS: Single
      - (D) TOPOLOGY: Linear
- 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

#### CACTGGCTCA GGGAAATAAC CC 22

55 (2) INFORMATION FOR SEQ ID NO:6:

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(i) SEQUENCE CHARACTERISTICS:
           (A) LENGTH: 22 base pairs
           (B) TYPE: Nucleic Acid
           (C) STRANDEDNESS: Single
           (D) TOPOLOGY: Linear
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      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
     GGAGAGCTGG GAAGGTGTGC AC 22
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     (2) INFORMATION FOR SEQ ID NO:7:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 35 base pairs
15
            (B) TYPE: Nucleic Acid
            (C) STRANDEDNESS: Single
            (D) TOPOLOGY: Linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:
20
      ACAAACGCGT ACGCTGACAT CGTCATGACC CAGTC 35
     (2) INFORMATION FOR SEQ ID NO:8:
25
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 35 base pairs
            (B) TYPE: Nucleic Acid
            (C) STRANDEDNESS: Single
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            (D) TOPOLOGY: Linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
35
      ACAAACGCGT ACGCTGATAT TGTCATGACT CAGTC 35
     (2) INFORMATION FOR SEQ ID NO:9:
         (i) SEQUENCE CHARACTERISTICS:
40
             (A) LENGTH: 35 base pairs
             (B) TYPE: Nucleic Acid
             (C) STRANDEDNESS: Single
             (D) TOPOLOGY: Linear
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        (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
       ACAAACGCGT ACGCTGACAT CGTCATGACA CAGTC 35
 50
      (2) INFORMATION FOR SEQ ID NO:10:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 37 base pairs
             (B) TYPE: Nucleic Acid
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             (C) STRANDEDNESS: Single
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(D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: 5 GCTCTTCGAA TGGTGGGAAG ATGGATACAG TTGGTGC 37 (2) INFORMATION FOR SEQ ID NO:11: (i) SEQUENCE CHARACTERISTICS: 10 (A) LENGTH: 39 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11: CGATGGGCCC GGATAGACCG ATGGGGCTGT TGTTTTGGC 39 20 (2) INFORMATION FOR SEQ ID NO:12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs 25 (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12: 30 CGATGGGCCC GGATAGACTG ATGGGGCTGT CGTTTTGGC 39 (2) INFORMATION FOR SEQ ID NO:13: 35 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single 40 (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13: CGATGGGCCC GGATAGACGG ATGGGGCTGT TGTTTTGGC 39 45 (2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 39 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
  - (D) TOPOLOGY: Linear
- 55 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

# CGATGGGCCC GGATAGACAG ATGGGGCTGT TGTTTTGGC 39

(2) INFORMATION FOR SEQ ID NO:15:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 39 base pairs
    - (B) TYPE: Nucleic Acid
    - (C) STRANDEDNESS: Single

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- (D) TOPOLOGY: Linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:
- 15 CGATGGGCCC GGATAGACCG ATGGGGCTGT TGTTTTGGC 39
  - (2) INFORMATION FOR SEQ ID NO:16:
    - (i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 39 base pairs
- (B) TYPE: Nucleic Acid
- (C) STRANDEDNESS: Single
- (D) TOPOLOGY: Linear
- 25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:
  - CGATGGGCCC GGATAGACTG ATGGGGCTGT TGTTTTGGC 39
- 30 (2) INFORMATION FOR SEQ ID NO:17:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 39 base pairs
    - (B) TYPE: Nucleic Acid
- 35 (C) STRANDEDNESS: Single
  - (D) TOPOLOGY: Linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

CGATGGGCCC GGATAGACAG ATGGGGCTGT TGTTTTGGC 39

- (2) INFORMATION FOR SEQ ID NO:18:
- 45 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 39 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
  - (D) TOPOLOGY: Linear

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- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:
- CGATGGGCCC GGATAGACGG ATGGGGCTGT TGTTTTGGC 39

55 (2) INFORMATION FOR SEQ ID NO:19:

(i) SEQUENCE CHARACTERISTICS:

5	<ul><li>(A) LENGTH: 369 base pairs</li><li>(B) TYPE: Nucleic Acid</li><li>(C) STRANDEDNESS: Double</li><li>(D) TOPOLOGY: Linear</li></ul>
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:
10	GACATTGTCA TGACACAGTC TCAAAAATTC ATGTCCACAT CAGTAGGAGA 50
	CAGGGTCAGC GTCACCTGCA AGGCCAGTCA GAATGTGGGT ACTAATGTAG 100
15	CCTGGTATCA ACAGAAACCA GGGCAATCTC CTAAAGCACT GATTTACTCG 150
	TCATCCTACC GGTACAGTGG AGTCCCTGAT CGCTTCACAG GCAGTGGATC 200
20	TGGGACAGAT TTCACTCTCA CCATCAGCCA TGTGCAGTCT GAAGACTTGG 250
	CAGACTATTT CTGTCAGCAA TATAACATCT ATCCTCTCAC GTTCGGTCCT 300
	GGGACCAAGC TGGAGTTGAA ACGGGCTGAT GCTGCACCAC CAACTGTATC 350
25	CATCTTCCCA CCATTCGAA 369
	(2) INFORMATION FOR SEQ ID NO:20:
30	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 123 amino acids</li><li>(B) TYPE: Amino Acid</li><li>(D) TOPOLOGY: Linear</li></ul>
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:
,,	Asp Ile Val Met Thr Gln Ser Gln Lys Phe Met Ser Thr Ser Val
40	Gly Asp Arg Val Ser Val Thr Cys Lys Ala Ser Gln Asn Val Gly 20 25 30
	Thr Asn Val Ala Trp Tyr Gln Gln Lys Pro Gly Gln Ser Pro Lys 35 40 49
45	Ala Leu Ile Tyr Ser Ser Ser Tyr Arg Tyr Ser Gly Val Pro Ass
50	Arg Phe Thr Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile 65 70 75
	Ser His Val Gln Ser Glu Asp Leu Ala Asp Tyr Phe Cys Gln Glr 80 85 90

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Tyr Asn Ile Tyr Pro Leu Thr Phe Gly Pro Gly Thr Lys Leu Glu

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ti ti tika kaja ja ala algada di kataja. Etti darri teda eta arka algada kataja ala da da da arka<u>ka katikarrirr</u>

Leu Lys Arg Ala Asp Ala Ala Pro Pro Thr Val Ser Ile Phe Pro 110 115 120

Pro Phe Glu 123

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- (2) INFORMATION FOR SEQ ID NO:21:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 417 base pairs
    - (B) TYPE: Nucleic Acid
    - (C) STRANDEDNESS: Double
    - (D) TOPOLOGY: Linear
- 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

TTCTATTGCT ACAAACGCGT ACGCTGAGGT GCAGCTGGTG GAGTCTGGGG 50

20 GAGGCTTAGT GCCGCCTGGA GGGTCCCTGA AACTCTCCTG TGCAGCCTCT 100

GGATTCATAT TCAGTAGTTA TGGCATGTCT TGGGTTCGCC AGACTCCAGG 150

CAAGAGCCTG GAGTTGGTCG CAACCATTAA TAATAATGGT GATAGCACCT 200

25
ATTATCCAGA CAGTGTGAAG GGCCGATTCA CCATCTCCCG AGACAATGCC 250

AAGAACACCC TGTACCTGCA AATGAGCAGT CTGAAGTCTG AGGACACAGC 300

ACTGGGGCCA AGGGACTCTG GTCACTGTCT CTGCAGCCAA AACAACAGCC 400

CCATCTGTCT ATCCGGG 417

(2) INFORMATION FOR SEQ ID NO:22:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 130 amino acids
- (B) TYPE: Amino Acid
  - (D) TOPOLOGY: Linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:
- 45 Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Pro Pro Gly
  1 5 10 15

Gly Ser Leu Lys Leu Ser Cys Ala Ala Ser Gly Phe Ile Phe Ser 20 25 30

Ser Tyr Gly Met Ser Trp Val Arg Gln Thr Pro Gly Lys Ser Leu
35 40 45

Glu Leu Val Ala Thr Ile Asn Asn Gly Asp Ser Thr Tyr Tyr
50 55 60

Pro Asp Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala 65 70 Lys Asn Thr Leu Tyr Leu Gln Met Ser Ser Leu Lys Ser Glu Asp 5 Thr Ala Met Phe Tyr Cys Ala Arg Ala Leu Ile Ser Ser Ala Thr 100 Trp Phe Gly Tyr Trp Gly Gln Gly Thr Leu Val Thr Val Ser Ala 10 Ala Lys Thr Thr Ala Pro Ser Val Tyr Pro 130 125 15 (2) INFORMATION FOR SEQ ID NO:23: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 base pairs 20 (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23: 25 ACAAACGCGT ACGCTGATAT CGTCATGACA G 31 (2) INFORMATION FOR SEQ ID NO:24: 30 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single 35 (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24: 40 GCAGCATCAG CTCTTCGAAG CTCCAGCTTG G 31 (2) INFORMATION FOR SEQ ID NO:25: (i) SEQUENCE CHARACTERISTICS: 45 (A) LENGTH: 21 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear 50 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25: CCACTAGTAC GCAAGTTCAC G 21 (2) INFORMATION FOR SEQ ID NO:26: 55

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 33 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
- 5 (D) TOPOLOGY: Linear

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:
- 10 GATGGGCCCT TGGTGGAGGC TGCAGAGACA GTG 33
  - (2) INFORMATION FOR SEQ ID NO:27:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 714 base pairs
      - (B) TYPE: Nucleic Acid
      - (C) STRANDEDNESS: Double
      - (D) TOPOLOGY: Linear
- 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:
  - ATGAAGAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT 50
- 25 TGCTACAAAC GCGTACGCTG ATATCGTCAT GACACAGTCT CAAAAATTCA 100
  - TGTCCACATC AGTAGGAGAC AGGGTCAGCG TCACCTGCAA GGCCAGTCAG 150
- AATGTGGGTA CTAATGTAGC CTGGTATCAA CAGAAACCAG GGCAATCTCC 200
- 30 TAAAGCACTG ATTTACTCGT CATCCTACCG GTACAGTGGA GTCCCTGATC 250
  - GCTTCACAGG CAGTGGATCT GGGACAGATT TCACTCTCAC CATCAGCCAT 300
- 35 GTGCAGTCTG AAGACTTGGC AGACTATTTC TGTCAGCAAT ATAACATCTA 350
  - TCCTCTCACG TTCGGTCCTG GGACCAAGCT GGAGCTTCGA AGAGCTGTGG 400
  - CTGCACCATC TGTCTTCATC TTCCCGCCAT CTGATGAGCA GTTGAAATCT 450
- 40 GGAACTGCTT CTGTTGTGTG CCTGCTGAAT AACTTCTATC CCAGAGAGGC 500
  - CAAAGTACAG TGGAAGGTGG ATAACGCCCT CCAATCGGGT AACTCCCAGG 550
- 45 AGAGTGTCAC AGAGCAGGAC AGCAAGGACA GCACCTACAG CCTCAGCAGC 600
  - ACCCTGACGC TGAGCAAAGC AGACTACGAG AAACACAAAG TCTACGCCTG 650
  - CGAAGTCACC CATCAGGGCC TGAGCTCGCC CGTCACAAAG AGCTTCAACA 700
- 50 GGGGAGAGTG TTAA 714
  - (2) INFORMATION FOR SEQ ID NO:28:
- 55 (i) SEQUENCE CHARACTERISTICS:
  (A) LENGTH: 237 amino acids

(B) TYPE: Amino Acid(D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

5 Met Lys Lys Asn Ile Ala Phe Leu Leu Ala Ser Met Phe Val Phe Ser Ile Ala Thr Asn Ala Tyr Ala Asp Ile Val Met Thr Gln Ser 10 20 Gln Lys Phe Met Ser Thr Ser Val Gly Asp Arg Val Ser Val Thr Cys Lys Ala Ser Gln Asn Val Gly Thr Asn Val Ala Trp Tyr Gln 15 Gln Lys Pro Gly Gln Ser Pro Lys Ala Leu Ile Tyr Ser Ser Ser 20 Tyr Arg Tyr Ser Gly Val Pro Asp Arg Phe Thr Gly Ser Gly Ser 85 Gly Thr Asp Phe Thr Leu Thr Ile Ser His Val Gln Ser Glu Asp 25 95 100 Leu Ala Asp Tyr Phe Cys Gln Gln Tyr Asn Ile Tyr Pro Leu Thr Phe Gly Pro Gly Thr Lys Leu Glu Leu Arg Arg Ala Val Ala Ala 30 125 130 Pro Ser Val Phe Ile Phe Pro Pro Ser Asp Glu Gln Leu Lys Ser 35 Gly Thr Ala Ser Val Val Cys Leu Leu Asn Asn Phe Tyr Pro Arg 160 155 Glu Ala Lys Val Gln Trp Lys Val Asp Asn Ala Leu Gln Ser Gly 175 40 Asn Ser Gln Glu Ser Val Thr Glu Gln Asp Ser Lys Asp Ser Thr 185 190 Tyr Ser Leu Ser Ser Thr Leu Thr Leu Ser Lys Ala Asp Tyr Glu 45 200 Lys His Lys Val Tyr Ala Cys Glu Val Thr His Gln Gly Leu Ser 50 Ser Pro Val Thr Lys Ser Phe Asn Arg Gly Glu Cys 237 230 235

(2) INFORMATION FOR SEQ ID NO:29:

55

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 756 base pairs

- (B) TYPE: Nucleic Acid
- (C) STRANDEDNESS: Double
- (D) TOPOLOGY: Linear

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

- ATGAAAAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT 50 10 TGCTACAAAC GCGTACGCTG AGGTGCAGCT GGTGGAGTCT GGGGGAGGCT 100 TAGTGCCGCC TGGAGGGTCC CTGAAACTCT CCTGTGCAGC CTCTGGATTC 150 ATATTCAGTA GTTATGGCAT GTCTTGGGTT CGCCAGACTC CAGGCAAGAG 200 15 CCTGGAGTTG GTCGCAACCA TTAATAATAA TGGTGATAGC ACCTATTATC 250 CAGACAGTGT GAAGGGCCGA TTCACCATCT CCCGAGACAA TGCCAAGAAC 300 20 ACCCTGTACC TGCAAATGAG CAGTCTGAAG TCTGAGGACA CAGCCATGTT 350 TTACTGTGCA AGAGCCCTCA TTAGTTCGGC TACTTGGTTT GGTTACTGGG 400 GCCAAGGGAC TCTGGTCACT GTCTCTGCAG CCTCCACCAA GGGCCCATCG 450 25 GTCTTCCCCC TGGCACCCTC CTCCAAGAGC ACCTCTGGGG GCACAGCGGC 500 CCTGGGCTGC CTGGTCAAGG ACTACTTCCC CGAACCGGTG ACGGTGTCGT 550 30 GGAACTCAGG CGCCCTGACC AGCGGCGTGC ACACCTTCCC GGCTGTCCTA 600 CAGTCCTCAG GACTCTACTC CCTCAGCAGC GTGGTGACCG TGCCCTCCAG 650 CAGCTTGGGC ACCCAGACCT ACATCTGCAA CGTGAATCAC AAGCCCAGCA 700 35 ACACCAAGGT GGACAAGAAA GTTGAGCCCA AATCTTGTGA CAAAACTCAC 750 ACATGA 756
  - (2) INFORMATION FOR SEQ ID NO:30:

40

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 251 amino acids
  - (B) TYPE: Amino Acid
  - (D) TOPOLOGY: Linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:
- 50 Met Lys Lys Asn Ile Ala Phe Leu Leu Ala Ser Met Phe Val Phe 1 5 10 15
  - Ser Ile Ala Thr Asn Ala Tyr Ala Glu Val Gln Leu Val Glu Ser 20 25 30
- 55
  Gly Gly Gly Leu Val Pro Pro Gly Gly Ser Leu Lys Leu Ser Cys

					35					40					45
5	Ala	Ala	Ser	Gly	Phe 50	Ile	Phe	Ser	Ser	Туг 55	Gly	Met	Ser	Trp	Val 60
3	Arg	Gln	Thr	Pro	Gly 65	Lys	Ser	Leu	Glu	Leu 70	Val	Ala	Thr	Ile	Asn 75
10	Asn	Asn	Gly	Asp	Ser 80	Thr	Tyr	Tyr	Pro	Asp 85	Ser	Val	Lys	Gly	Arg 90
	Phe	Thr	Ile	Ser	Arg 95	Asp	Asn	Ala	Lys	Asn 100	Thr	Leu	Tyr	Leu	Gln 105
15	Met	Ser	Ser	Leu	Lys 110	Ser	Glu	Asp	Thr	Ala 115	Met	Phe	Tyr	Cys	Ala 120
20	Arg	Ala	Leu	Ile	Ser 125	Ser	Ala	Thr	Trp	Phe 130	Gly	Tyr	Trp	Gly	Gln 135
	Gly	Thr	Leu	Val	Thr 140	Val	Ser	Ala	Ala	Ser 145	Thr	Lys	Gly	Pro	Ser 150
25	Val	Phe	Pro	Leu	Ala 155	Pro	Ser	Ser	Lys	Ser 160	Thr	Ser	Gly	Gly	Thr 165
	Ala	Ala	Leu	Gly	Cys 170	Leu	Val	Lys	Asp	Tyr 175	Phe	Pro	Glu	Pro	Val 180
30	Thr	Val	Ser	Trp	Asn 185	Ser	Gly	Ala	Leu	Thr 190	Ser	Gly	Val	His	Thr 195
35	Phe	Pro	Ala	Val	Leu 200	Gln	Ser	Ser	Gly	Leu 205	Tyr	Ser	Leu	Ser	Ser 210
	Val	Val	Thr	Val	Pro 215	Ser	Ser	Ser	Leu	Gly 220	Thr	Gln	Thr	Tyr	Ile 225
40	Cys	Asn	Val	Asn	His 230	Lys	Pro	Ser	Asn	Thr 235	Lys	Val	Asp	Lys	Lys 240
	Val	Glu	Pro	Lys	Ser 245	Cys	Asp	Lys	Thr		Thr 251				
45	(2)	INFO	RMAT	ION I	FOR S	SEQ :	ID N	0:31	:						
	1.	i) e:	ישוזטפ	י שטע	CHAR	י בירי	7 T C T'	TCS ·							
	ν.				H: 2										
50					Nuc										
JU		((	<b>⊸</b> , ສີ	T KWN)	DEDNI	255:	PING	јте							

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

CAGTCCAACT GTTCAGGACG CC 22

55

(D) TOPOLOGY: Linear

	(2) INFORMATION FOR SEQ ID NO:32:
_	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs
5	(B) TYPE: Nucleic Acid
	(C) STRANDEDNESS: Single
	(D) TOPOLOGY: Linear
	(2) 2020000
0	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:
	GTGCTGCTCA TGCTGTAGGT GC 22
15	(2) INFORMATION FOR SEQ ID NO:33:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 23 base pairs
	(B) TYPE: Nucleic Acid
20	(C) STRANDEDNESS: Single
	(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:
25	·
	GAAGTTGATG TCTTGTGAGT GGC 23
	(2) INFORMATION FOR SEQ ID NO:34:
30	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 24 base pairs
	(B) TYPE: Nucleic Acid
	(C) STRANDEDNESS: Single
	(D) TOPOLOGY: Linear
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:
40	GCATCCTAGA GTCACCGAGG AGCC 24
40	(2) INFORMATION FOR SEQ ID NO:35:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 22 base pairs
45	(B) TYPE: Nucleic Acid
	(C) STRANDEDNESS: Single
	(D) TOPOLOGY: Linear
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:
50	
	CACTGGCTCA GGGAAATAAC CC 22
	(2) INFORMATION FOR SEQ ID NO:36:
55	(i) SPOUENCE CHARACTERISTICS:

276.3879 200.3879

(A) LENGTH: 22 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear 5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36: GGAGAGCTGG GAAGGTGTGC AC 22 10 (2) INFORMATION FOR SEQ ID NO:37: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 37 base pairs 15 (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:37: 20 CCAATGCATA CGCTGACATC GTGATGACCC AGACCCC 37 (2) INFORMATION FOR SEQ ID NO:38: 25 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 37 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single 30 (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38: 35 CCAATGCATA CGCTGATATT GTGATGACTC AGACTCC 37 (2) INFORMATION FOR SEQ ID NO:39: (i) SEQUENCE CHARACTERISTICS: 40 (A) LENGTH: 37 base pairs (B) TYPE: Nucleic Acid (C) STRANDEDNESS: Single (D) TOPOLOGY: Linear 45 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39: CCAATGCATA CGCTGACATC GTGATGACAC AGACACC 37 (2) INFORMATION FOR SEQ ID NO:40: 50 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 base pairs (B) TYPE: Nucleic Acid 55 (C) STRANDEDNESS: Single

(D) TOPOLOGY: Linear

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```
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:
     AGATGTCAAT TGCTCACTGG ATGGTGGGAA GATGG 35
5
    (2) INFORMATION FOR SEQ ID NO:41:
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 32 base pairs
10
            (B) TYPE: Nucleic Acid
            (C) STRANDEDNESS: Single
            (D) TOPOLOGY: Linear
      (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:
15
     CAAACGCGTA CGCTGAGATC CAGCTGCAGC AG 32
     (2) INFORMATION FOR SEQ ID NO:42:
20
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 32 base pairs
            (B) TYPE: Nucleic Acid
            (C) STRANDEDNESS: Single
25
            (D) TOPOLOGY: Linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:
30
      CAAACGCGTA CGCTGAGATT CAGCTCCAGC AG 32
     (2) INFORMATION FOR SEQ ID NO:43:
35
        (i) SEQUENCE CHARACTERISTICS:
            (A) LENGTH: 39 base pairs
            (B) TYPE: Nucleic Acid
            (C) STRANDEDNESS: Single
            (D) TOPOLOGY: Linear
40
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:
      CGATGGGCCC GGATAGACCG ATGGGGCTGT TGTTTTGGC 39
45
     (2) INFORMATION FOR SEQ ID NO:44:
         (i) SEQUENCE CHARACTERISTICS:
             (A) LENGTH: 39 base pairs
```

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:

(B) TYPE: Nucleic Acid

(C) STRANDEDNESS: Single
(D) TOPOLOGY: Linear

50

#### CGATGGGCCC GGATAGACTG ATGGGGCTGT TGTTTTGGC 39

```
(2) INFORMATION FOR SEQ ID NO:45:
```

- 5 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 39 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
  - (D) TOPOLOGY: Linear

10

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:

CGATGGGCCC GGATAGACAG ATGGGGCTGT TGTTTTGGC 39

15

20

- (2) INFORMATION FOR SEQ ID NO:46:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 39 base pairs
    - (B) TYPE: Nucleic Acid
      - (C) STRANDEDNESS: Double
      - (D) TOPOLOGY: Linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:

25

CGATGGGCCC GGATAGACGG ATGGGGCTGT TGTTTTGGC 39

(2) INFORMATION FOR SEQ ID NO:47:

30

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 391 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Double
- 35

- (D) TOPOLOGY: Linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:
- 40 GATATCGTGA TGACACAGAC ACCACTCTCC CTGCCTGTCA GTCTTGGAGA 50
  - TCAGGCCTCC ATCTCTTGCA GATCTAGTCA GAGCCTTGTA CACGGTATTG 100
- GAAACACCTA TTTACATTGG TACCTGCAGA AGCCAGGCCA GTCTCCAAAG 150
  - CTCCTGATCT ACAAAGTTTC CAACCGATTT TCTGGGGTCC CAGACAGGTT 200
    - CAGTGGCAGT GGATCAGGGA CAGATTTCAC ACTCAGGATC AGCAGAGTGG 250
- 50 AGGCTGAGGA TCTGGGACTT TATTTCTGCT CTCAAAGTAC ACATGTTCCG 300
  - CTCACGTTCG GTGCTGGGAC CAAGCTGGAG CTGAAACGGG CTGATGCTGC 350
    - ACCAACTGTA TCCATCTTCC CACCATCCAG TGAGCAATTG A 391
    - (2) INFORMATION FOR SEQ ID NO:48:

5	(1	( <i>P</i>	LE 3) TY	NGTH PE:	I: 13 Amin )GY:	1 am	ino id	acid	ls						
	(xi	) SE	QUEN	ICE I	ESCR	IPTI	ON:	SEQ	ID N	10:48	3:				
10	Asp 1	Ile	Val	Met	Thr 5	Gln	Thr	Pro	Leu	Ser 10	Leu	Pro	Val	Ser	Leu 15
	Gly	Asp	Gln	Ala	Ser 20	Ile	Ser	Cys	Arg	Ser 25	Ser	Gln	Ser	Leu	Val 30
15	His	Gly	Ile	Gly	Asn 35	Thr	Tyr	Leu	His	Trp 40	Tyr	Leu	Gln	Lys	Pro 45
20	Gly	Gln	Ser	Pro	Lys 50		Leu	Ile	Tyr	Lys 55	Val	Ser	Asn	Arg	Phe 60
20	Ser	Gly	Val	Pro	Asp 65	Arg	Phe	Ser	Gly	Ser 70	Gly	Ser	Gly	Thr	Asp 75
25	Phe	Thr	Leu	Arg	Ile 80	Ser	Arg	Val	Glu	Ala 85	Glu	Asp	Leu	Gly	Leu 90
,	Tyr	Phe	Cys	Ser	Gln 95	Ser	Thr	His	Val	Pro 100		Thr	Phe	Gly	Ala 105
30	Gly	Thr	Lys	Leu	Glu 110	Leu	Lys	Arg	Ala	Asp 115		Ala	Pro	Thr	Val 120
26	Ser	Ile	Phe	Pro	Pro 125	Ser	Ser	Glu	Gln		Lys 131				
35					FOR				:						
40		( (	A) L B) T C) S	ENGT YPE : TRAN	CHAR H: 4 'Nuc DEDN JOGY:	05 b leic ESS:	ase : Aci Dou	pair d	s						
45	(x	i) S	EQUE	NCE	DESC	RIPI	: NOI	SEC	D	NO : 4	19:				
	GAG	ATTC	CAGC	TGCA	GCAG	TC I	rggac	CTG	G CI	GATO	EAAGO	CTC	GGGG	CTTC	50
50	AGT	GAAG	ATA	TCCI	rgcaa	GG (	CTTCT	GGTT	TA T	CATI	rcag1	AG(	CACT	raca	100
					GCAGA										
					ATGGT										
55	GGC	CAC	ATTG	ACT	GTAG#	ACA (	CATC	rtcc)	AG C	ACAG	CCAA	GT	CAT	CTCA	250

。 1987年 - 19874 - 1987年 - 198

GCAGCCTGAC ATCTGATGAC TCTGCAGTCT ATTTCTGTGC AAGAGGGGAC 300
TATAGATACA ACGGCGACTG GTTTTTCGAT GTCTGGGGNG NAGGGACCAC 350
GGTCACCGTC TCCTCCGCCA AAACCGACAG CCCCATCGGT CTATCCGGGC 400
CCATC 405

(2) INFORMATION FOR SEQ ID NO:50:

10

5

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 135 amino acids
  - (B) TYPE: Amino Acid
  - (D) TOPOLOGY: Linear

15

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40

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

Glu Ile Gln Leu Gln Gln Ser Gly Pro Glu Leu Met Lys Pro Gly
1 5 10 15

20

Ala Ser Val Lys Ile Ser Cys Lys Ala Ser Gly Tyr Ser Phe Ser 20 25 30

Ser His Tyr Met His Trp Val Lys Gln Ser His Gly Lys Ser Leu 25 40 45

Glu Trp Ile Gly Tyr Ile Asp Pro Ser Asn Gly Glu Thr Thr Tyr
50 55 60

30 Asn Gln Lys Phe Lys Gly Lys Ala Thr Leu Thr Val Asp Thr Ser 65 70 75

Ser Ser Thr Ala Asn Val His Leu Ser Ser Leu Thr Ser Asp Asp 80 85 90

Ser Ala Val Tyr Phe Cys Ala Arg Gly Asp Tyr Arg Tyr Asn Gly
95 100 105

Asp Trp Phe Phe Asp Val Trp Gly Xaa Gly Thr Thr Val Thr Val

Ser Ser Ala Lys Thr Asp Ser Pro Ile Gly Leu Ser Gly Pro Ile 125 130 135

- 45 (2) INFORMATION FOR SEQ ID NO:51:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22 base pairs
    - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
    - (D) TOPOLOGY: Linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

55

50

CTTGGTGGAG GCGGAGGAGA CG 22

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	(2) INFORMATION FOR SEQ ID NO:52:
5	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 38 base pairs</li> <li>(B) TYPE: Nucleic Acid</li> <li>(C) STRANDEDNESS: Single</li> <li>(D) TOPOLOGY: Linear</li> </ul>
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:
	GAAACGGGCT GTTGCTGCAC CAACTGTATT CATCTTCC 38
15	(2) INFORMATION FOR SEQ ID NO:53:
20	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 31 base pairs</li> <li>(B) TYPE: Nucleic Acid</li> <li>(C) STRANDEDNESS: Single</li> <li>(D) TOPOLOGY: Linear</li> </ul>
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:
	(X1) SEQUENCE DESCRIPTION: SEQ 15 No. 33.
25	GTCACCGTCT CCTCCGCCTC CACCAAGGGC C 31
	(2) INFORMATION FOR SEQ ID NO:54:
30	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 22 base pairs</li> <li>(B) TYPE: Nucleic Acid</li> <li>(C) STRANDEDNESS: Single</li> <li>(D) TOPOLOGY: Linear</li> </ul>
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:
	CTTGGTGGAG GCGGAGGAGA CG 22
40	(2) INFORMATION FOR SEQ ID NO:55:
45	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 729 base pairs</li> <li>(B) TYPE: Nucleic Acid</li> <li>(C) STRANDEDNESS: Double</li> <li>(D) TOPOLOGY: Linear</li> </ul>
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:
	ATGAAGAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT 50
	TGCTACAAAT GCATACGCTG ATATCGTGAT GACACAGACA CCACTCTCCC 100
55	TGCCTGTCAG TCTTGGAGAT CAGGCCTCCA TCTCTTGCAG ATCTAGTCAG 150

	AGCCTTGTAC	ACGGTATTGG	AAACACCTAT	TTACATTGGT	ACCTGCAGAA	200
5	GCCAGGCCAG	TCTCCAAAGC	TCCTGATCTA	CAAAGTTTCC	AACCGATTTT	250
J	CTGGGGTCCC	AGACAGGTTC	AGTGGCAGTG	GATCAGGGAC	AGATTTCACA	300
	CTCAGGATCA	GCAGAGTGGA	GGCTGAGGAT	CTGGGACTTT	ATTTCTGCTC	350
10	TCAAAGTACA	CATGTTCCGC	TCACGTTCGG	TGCTGGGACC	AAGCTGGAGC	400
	TGAAACGGGC	TGTTGCTGCA	CCAACTGTAT	TCATCTTCCC	ACCATCCAGT	450
15	GAGCAATTGA	AATCTGGAAC	TGCCTCTGTT	GTGTGCCTGC	TGAATAACTT	500
	CTATCCCAGA	GAGGCCAAAG	TACAGTGGAA	GGTGGATAAC	GCCCTCCAAT	550
	CGGGTAACTC	CCAGGAGAGT	GTCACAGAGC	AGGACAGCAA	GGACAGCACC	600
20	TACAGCCTCA	GCAGCACCCT	GACGCTGAGC	AAAGCAGACT	ACGAGAAACA	650
	CAAAGTCTAC	GCCTGCGAAG	TCACCCATCA	GGGCCTGAGC	TCGCCCGTCA	700
25	CAAAGAGCTT	CAACAGGGGA	GAGTGTTAA	729		
	(2) INFORMAT	TION FOR SEC	D ID NO:56:			

- - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 242 amino acids
  - (B) TYPE: Amino Acid
    - (D) TOPOLOGY: Linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:
- Met Lys Lys Asn Ile Ala Phe Leu Leu Ala Ser Met Phe Val Phe 35
- Ser Ile Ala Thr Asn Ala Tyr Ala Asp Ile Val Met Thr Gln Thr 40
  - Pro Leu Ser Leu Pro Val Ser Leu Gly Asp Gln Ala Ser Ile Ser 40 35
- Cys Arg Ser Ser Gln Ser Leu Val His Gly Ile Gly Asn Thr Tyr 45
  - Leu His Trp Tyr Leu Gln Lys Pro Gly Gln Ser Pro Lys Leu Leu 65
- 50 Ile Tyr Lys Val Ser Asn Arg Phe Ser Gly Val Pro Asp Arg Phe
  - Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Arg Ile Ser Arg 95 100
- 55 Val Glu Ala Glu Asp Leu Gly Leu Tyr Phe Cys Ser Gln Ser Thr

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	110 115 120
	His Val Pro Leu Thr Phe Gly Ala Gly Thr Lys Leu Glu Leu Lys 125 130 135
5	Arg Ala Val Ala Ala Pro Thr Val Phe Ile Phe Pro Pro Ser Ser 140
10	Glu Gln Leu Lys Ser Gly Thr Ala Ser Val Val Cys Leu Leu Asn 155 160 165
	Asn Phe Tyr Pro Arg Glu Ala Lys Val Gln Trp Lys Val Asp Asn 170 175 180
15	Ala Leu Gln Ser Gly Asn Ser Gln Glu Ser Val Thr Glu Gln Asp 185 190 195
	Ser Lys Asp Ser Thr Tyr Ser Leu Ser Ser Thr Leu Thr Leu Ser 200 205 210
20	Lys Ala Asp Tyr Glu Lys His Lys Val Tyr Ala Cys Glu Val Thr 215 220 225
25	His Gln Gly Leu Ser Ser Pro Val Thr Lys Ser Phe Asn Arg Gly 230 235 240
	Glu Cys 242
30	(2) INFORMATION FOR SEQ ID NO:57:
35	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 762 base pairs</li> <li>(B) TYPE: Nucleic Acid</li> <li>(C) STRANDEDNESS: Double</li> <li>(D) TOPOLOGY: Linear</li> </ul>
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:
40	ATGAAAAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT 50
	TGCTACAAAC GCGTACGCTG AGATTCAGCT GCAGCAGTCT GGACCTGAGC 100
45	TGATGAAGCC TGGGGCTTCA GTGAAGATAT CCTGCAAGGC TTCTGGTTAT 150
	TCATTCAGTA GCCACTACAT GCACTGGGTG AAGCAGAGCC ATGGAAAGAG 200
	CCTTGAGTGG ATTGGCTACA TTGATCCTTC CAATGGTGAA ACTACTTACA 250
50	ACCAGAAATT CAAGGGCAAG GCCACATTGA CTGTAGACAC ATCTTCCAGC 300
	ACAGCCAACG TGCATCTCAG CAGCCTGACA TCTGATGACT CTGCAGTCTA 350
55	TTTCTGTGCA AGAGGGGACT ATAGATACAA CGGCGACTGG TTTTTCGATG 400

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	TCTG	GGGG	CGC .	AGGG	ACCA	CG G	TCAC	CGTC	T CC	TCCG	CCTC	CAC	CAAG	GGC	450
	CCAT	CGG:	TCT '	TCCC	CCTG	GC A	.CCCT	CCTC	C AA	GAGC	ACCT	CTG	GGGG	CAC	500
5	AGCG	GCC	CTG.	GGCT	GCCT	GG T	CAAG	GACT	A CT	TCCC	CGAA	CCG	GTGA	CGG	550
	TGTC	GTG	GAA (	CTCA	GGCG	cc c	TGAC	CAGC	G GC	GTGC	ACAC	CTT	CCCG	GCT	600
10	GTCC	TAC	AGT (	CCTC	AGGA	CT C	TACT	CCCT	C AG	CAGC	GTGG	TGA	CCGT	GCC	650
10	CTCC	:AGC	AGC '	TTGG	GCAC	CC A	GACC	TACA	r cr	GCAA	CGTG	AAT	CACA	AGC	700
	CCAG	CAAC	CAC	CAAG	GTGG	AC A	AGAA	AGTT	G AG	CCCA	AATC	TTG	TGAC	AAA	750
15	ACTO	:ACA(	CAT (	GA 7	52										
	(2) I							•	•						
20	(1	( <i>I</i>	A) L1 3) T	ENGTI YPE :	CHARA H: 29 Amir OGY:	53 a	mino cid		is						
25	(xi	) SE	EQUE	NCE I	DESCI	RIPT	ION:	SEQ	ID I	NO : 5	B:				
	Met 1	Lys	Lys	Asn	Ile 5	Ala	Phe	Leu	Leu	Ala 10	Ser	Met	Phe	Val	Phe 15
30	Ser	Ile	Ala	Thr	Asn 20	Ala	Tyr	Ala	Glu	Ile 25	Gln	Leu	Gln	Gln	Ser 30
	Gly	Pro	Glu	Leu	Met 35	Lys	Pro	Gly	Ala	Ser 40	Val	Lys	Ile	Ser	Cys 45
35	Lys .	Ala	Ser	Gly	Tyr 50	Ser	Phe	Ser	Ser	His 55	Tyr	Met	His	Trp	Val 60
40	Lys	Gln	Ser	His	Gly 65	Lys	Ser	Leu	Glu	Trp 70	Ile	Gly	Tyr	Ile	Asp 75
.0	Pro	Ser	Asn	Gly	Glu 80	Thr	Thr	Tyr	Asn	Gln 85	Lys	Phe	Lys	Gly	Lys 90
45	Ala	Thr	Leu	Thr	Val 95	Asp	Thr	Ser	Ser	Ser 100	Thr	Ala	Asn	Val	His 105
	Leu	Ser	Ser	Leu	Thr 110	Ser	Asp	Asp	Ser	Ala 115	Val	Tyr	Phe	Cys	Ala 120
50	Arg (	Gly	Asp	Tyr	Arg 125	Tyr	Asn	Gly	Asp	Trp 130	Phe	Phe	Asp	Val	Trp 135
55	Gly 2	Ala	Gly	Thr	Thr 140	Val	Thr	Val	Ser	Ser 145	Ala	Ser	Thr	Lys	Gly 150
<i></i>	Pro s	Ser	Val	Phe	Pro	Leu	Ala	Pro	Ser	Ser	Lys	Ser	Thr	Ser	Gly

					155					160					165
	Gly.	Thr	Ala	Ala	Leu 170	Gly	Cys	Leu	Val	Lys 175	Ąsp	Tyr	Phe	Pro	Glu 180
5	Pro	Val	Thr	Val	Ser 185	Trp	Asn	Ser	Gly	Ala 190	Leu	Thr	Ser	Gly	Val 195
0	His	Thr	Phe	Pro	Ala 200	Val	Leu	Gln	Ser	Ser 205	Gly	Leu	Tyr	Ser	Leu 210
	Ser	Ser	Val	Val	Thr 215	Val	Pro	Ser	Ser	Ser 220	Leu	Gly	Thr	Gln	Thr 225
15	Tyr	Ile	Cys	Asn	Val 230	Asn	His	Lys	Pro	Ser 235	Asn	Thr	Lys	Val	Asp 240
	Lys	Lys	Val	Glu	Pro 245	Lys	Ser	Cys	Asp	Lys 250	Thr	His	Thr 253	•	
20	(2)	INFO	RMAT	ION	FOR S	SEQ :	ID NO	5:59	:						•
25	€.	(	A) L B) T	ENGT YPE :	CHARI H: 1: Amii OGY:	14 an	mino cid		ds						
	(x	i) S	EQUE	NCE	DESC	RIPT	: NOI	SEQ	ID	NO:5	9:				
30	Asp 1		val	Met	Thr 5	Gln	Thr	Pro	Leu	Ser 10		Pro	Val	Ser	Leu 15
25	Gly	Asp	Gln	Ala	Ser 20	Ile	Ser	Cys	Arg	Ser 25		Glr	Ser	Leu	Val 30
35	His	Gly	/ Ile	e Gly	Asn 35		туг	Lev	ı His	Trp		Lev	ı Glr	Lys	Pro 45
40	Gly	Gl:	ı Sei	Pro	Lys 50		Leu	Ile	≥ Ту	7 Tyr 55		s Val	l Ser	Asr	Arg 60
	Ph∈	e Se	r Gly	y Val	l Pro		Arg	Phe	e Se:	r Asp 70		r Gly	y Sei	Gly	/ Thr 75
45	Asp	) Ph	e Th:	r Lei	a Arg		e Ser	Ar	g Va	1 Gl:		a Gl	u Asj	p Lev	ı Gly 90
50	Lev	з Ту	r Ph	е Су	s Sei 99		ı Sei	Th	r Hi	s Va 10		o Le	u Th	r Pho	e Gly 109
50	Ala	a Gl	уTh	r Ly	s Lei 110		ı Lev	ı Ly	s Ar 11						
55	(2)				FOR										

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(A) LENGTH: 114 amino acids (B) TYPE: Amino Acid (D) TOPOLOGY: Linear 5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60: Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val Gly Asp Arg Val Thr Ile Thr Cys Arg Ser Ser Gln Ser Leu Val 10 His Gly Ile Gly Asn Thr Tyr Leu His Trp Tyr Gln Gln Lys Pro 15 Gly Lys Ala Pro Lys Leu Leu Ile Tyr Tyr Lys Val Ser Asn Arg Phe Ser Gly Val Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr 20 Asp Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro Glu Asp Phe Ala 80 Thr Tyr Tyr Cys Ser Gln Ser Thr His Val Pro Leu Thr Phe Gly 25 100 Gln Gly Thr Lys Val Glu Ile Lys Arg 110 30 (2) INFORMATION FOR SEQ ID NO:61: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 109 amino acids 35 (B) TYPE: Amino Acid (D) TOPOLOGY: Linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:61: 40 Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val 5 1 Gly Asp Arg Val Thr Ile Thr Cys Arg Ala Ser Lys Thr Ile Ser 45 Lys Tyr Leu Ala Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu Ile Tyr Tyr Ser Gly Ser Thr Leu Glu Ser Gly Val Pro 50 Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile Ser Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln

	Gln Hi	s Asn	Glu	Tyr 95	Pro	Leu	Thr	Phe	Gly 100	Gln	Gly	Thr	Lys	Val 105
5	Glu Il	e Lys	Arg 109					•						
	(2) INF	ORMAT	ON F	OR S	EQ 1	D NC	:62:							
10		SEQUEI (A) LI (B) TI (D) TO	ENGTH YPE :	I: 11 Amir	17 an	nino cid		ls						
15	(xi)	SEQUE	NCE I	ESCF	RIPT	ON:	SEQ	ID 1	10:62	2:				
	Glu Il 1	e Gln	Leu	Gln 5	Gln	Ser	Gly	Pro	Glu 10	Leu	Met	Lys	Pro	Gly 15
20	Ala Se	r Val	Lys	Ile 20	Ser	Cys	Lys	Aļa	Ser 25	Gly	Tyr	Ser	Phe	Ser 30
0.5	Ser Hi	s Tyr	Met	His 35	Trp	Val	Lys	Gln	Ser 40	His	Gly	Lys	Ser	Leu 45
25	Glu Tr	p Ile	Gly	Туг 50	Ile	Asp	Pro	Ser	Asn 55	Gly	Glu	Thr	Thr	Tyr 60
30	Asn Gl	n Lys	Phe	Lys 65	Gly	Lys	Ala	Thr	Leu 70	Thr	Val	Asp	Thr	Ser 75
	Ser Se	r Thr	Ala	Asn 80		His	Leu	Ser	Ser 85	Leu	Thr	Ser	Asp	Asp 90
35	Ser Al	a Val	Tyr	Phe 95		Ala	Ala	Arg	Gly 100		туг	Arg	Tyr	Asn 105
	Gly. As	p Trp	Phe	Phe 110		Val	Trp	Gly	Ala 115		Thr 117			
40	(2) INI	FORMAT	NOI	FOR	SEQ	ID N	0:63	:						
45	(i)	SEQUE (A) I (B) I	ENGT	H: l Ami	.17 a	mino cid								
	(xi)	SEQUI	ENCE	DESC	RIPT	:NOI	SEÇ	O ID	NO : 6	3:				
50	Glu V	al Glı	n Lev	val		ı Ser	Gly	y Gly	/ Gly		ı Val	Glr	ı Pro	Gly 15
	Gly S	er Le	ı Arg	Lev 20		c Cys	s Ala	a Ala	a Ser 25		у Туз	r Sei	r Phe	e Ser 30
55	Ser H	is Ty:	r Met	: His	s Tr	y Val	l Arg	g Glı	n Ala	a Pro	o Gly	y Lys	s Gly	y Leu

					35					40					45
5	Glu	Trp	Val	Gly	Туг 50	Ile	Asp	Pro	Ser	Asn 55	Gly	Glu	Thr	Thr	Ту1 60
	Asn	Gln	Lys	Phe	Lys 65	Gly	Arg	Phe	Thr	Ile 70	Ser	Arg	Asp	Asn	Ser 75
10	Lys	Asn	Thr	Leu	Tyr 80	Leu	Gln	Met	Asn	Ser 85	Leu	Arg	Ala	Glu	Asp 90
	Thr	Ala	Val	Tyr	Tyr 95	Cys	Ala	Ala	Arg	Gly 100	Asp	Tyr	Arg	Tyr	Asr 105
15	Gly	Asp	Trp	Phe	Phe 110	Asp	Val	Trp	Gly	Gln 115	Gly	Thr 117			
	(2)	INFO	RMAT:	ION I	FOR S	SEQ :	ID NO	0:64	:						
20	(	() (1	EQUEI A) Li B) Ti	ENGTI YPE :	H: 1: Amiı	16 ar	mino cid	_	ls						٠
25	. (x	i) SI	EQUE	NCE 1	DESCI	RIPT:	ION:	SEQ	ID 1	10:64	<b>l</b> :				
	Glu 1	Val	Gln	Leu	Val 5	Glu	Ser	Gly	Gly	Gly 10	Leu	Val	Gln	Pro	Gly 15
30	Gly	Ser	Leu	Arg	Leu 20	Ser	Cys	Ala	Ala	Ser '25	Gly	Phe	Ser	Phe	Thr 30
35	Gly	His	Trp	Met	Asn 35	Trp	Val	Arg	Gln	Ala 40	Pro	Gly	Lys	Gly	Let 45
	Glu	Trp	Val	Gly	Met 50	Ile	His	Pro	Ser	Asp 55	Ser	Glu	Thr	Arg	Ту: 60
40	Ala	Asp	Ser	Val	Lys 65	Gly	Arg	Phe	Thr	Ile 70	Ser	Arg	Asp	Asn	Sex 75
	Lys	Asn	Thr	Leu	Tyr 80	Leu	Gln	Met	Asn	Ser 85	Leu	Arg	Ala	Glu	Asp 90
45	Thr	Ala	Val	Tyr	Tyr 95	Cys	Ala	Ala	Arg	Gly 100	Ile	Tyr	Phe	Tyr	Gl <sub>3</sub>
50			Tyr RMAT:		110						Thr 116				
55	(	() ()	EQUE A) L B) T D) T	ENGT YPE :	H: 2	42 at no A	mino cid		ds						

and the state of the field of the state of the

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:65:

5	Met 1	Lys	Lys	Asn	Ile 5	Ala	Phe	Leu	Leu	Ala 10	Ser	Met	Phe	Val	Phe 15
	Ser	Ile	Ala	Thr	Asn 20	Ala	Tyr	Ala	Asp	Ile 25	Gln	Met	Thr	Gln	Ser 30
10	Pro	Ser	Ser	Leu	Ser 35	Ala	Ser	Val	Gly	Asp 40	Arg	Val	Thr	Ile	Thr 45
	Cys	Arg	Ser	Ser	Gln 50	Ser	Leu	Val	His	Gly 55	Ile	Gly	Asn	Thr	Tyr 60
15	Leu	His	Trp	Tyr	Gln 65	Gln	Lys	Pro	Gly	Lys 70	Ala	Pro	Lys	Leu	Leu 75
20	Ile	Tyr	Lys	Val	Ser 80	Asn	Arg	Phe	Ser	Gly 85		Pro	Ser	Arg	Phe 90
	Ser	Gly	Ser	Gly	Ser 95	Gly	Thr	Asp	Phe	Thr 100	Leu	Thr	Ile	Ser	Ser 105
25	Leu	Gln	Pro	Glu	Asp 110	Phe	Ala	Thr	Tyr	Tyr 115	Cys	Ser	Gln	Ser	Thr 120
	His	Val	Pro	Leu	Thr 125	Phe	Gly	Gln	Gly	Thr 130	Lys	Val	Glu	Ile	Lys 135
30	Arg	Thr	Val	Ala	Ala 140	Pro	Ser	Val	Phe	Ile 145		Pro	Pro	Ser	Asp 150
35	Glu	Gln	Leu	Lys	Ser 155	Gly	Thr	Ala	Ser	Val 160		Cys	Leu	Leu	Asn 165
	Asn	Phe	Tyr	Pro	Arg 170	Glu	Ala	Lys	Val	Gln 175		Lys	Val	Asp	Asn 180
40	Ala	Leu	Gln	Ser	Gly 185		Ser	Gln	Glu	Ser 190		Thr	Glu	Gln	Asp 195
	Ser	Lys	Asp	Ser	Thr 200		Ser	Leu	Ser	Ser 205		Leu	Thr	Leu	Ser 210
<b>45</b>	Lys	: Ala	Asp	Tyr	Glu 215		His	Lys	. Val	Tyr 220		Cys	: Glu	Val	Thr 225
50	His	Glr	Gly	Leu	Ser 230		Pro	Val	. Thr	Lys 235		Phe	a Asn	Arg	Gly 240
	Glu	ı Cys 242													
									_						

55 (2) INFORMATION FOR SEQ ID NO:66:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 253 amino acids
- (B) TYPE: Amino Acid
- (D) TOPOLOGY: Linear

5	(xi) SE	QUENCE I	ESCRIPTI	ON: SEQ	ID NO:6	6:		
10	Met Lys	Lys Asn	Ile Ala 5	Phe Leu	Leu Ala 10		Phe Val	Phe 15
10	Ser Ile	Ala Thr	Asn Ala 20	Tyr Ala	Glu Val 25	Gln Leu	Val Gln	Ser 30
15	Gly Gly	Gly Leu	Val Gln 35	Pro Gly	Gly Ser	Leu Arg	Leu Ser	Cys 45
	Ala Ala	Ser Gly	Tyr Ser 50	Phe Ser	Ser His 55	Tyr Met	His Trp	Val 60
20	Arg Gln	Ala Pro	Gly Lys 65	Gly Let	Glu Trp 70	Val Gly	Tyr Ile	Asp 75
25	Pro Ser	Asn Gly	Glu Thr 80	Thr Ty	Asn Gln 85	Lys Phe	Lys Gly	Arg 90
	Phe Thr	Leu Ser	Arg Asp 95	Asn Ser	Lys Asn 100		Tyr Leu	Gln 105
30	Met Asn	Ser Leu	Arg Ala 110	Glu Asp	Thr Ala		Tyr Cys	Ala 120
	Arg Gly	Asp Tyr	Arg Tyr 125	Asn Gly	Asp Trp 130		Asp Val	Trp 135
35	Gly Gln	Gly Thr	Leu Val 140	Thr Va	l Ser Ser 145		Thr Lys	Gly 150
40	Pro Ser	Val Phe	Pro Leu 155	Ala Pro	Ser Ser 160		Thr Ser	Gly 165
	Gly Thr	Ala Ala	Leu Gly 170	Cys Let	u Val Lys 175		Phe Pro	Glu 180
45	Pro Val	Thr Val	Ser Trp 185	Asn Se	r Gly Ala 190		Ser Gly	Val 195
	His Thr	Phe Pro	Ala Val 200	Leu Gl	n Ser Ser 205		Tyr Ser	Leu 210
50	Ser Ser	Val Val	Thr Val 215	Pro Se	r Ser Ser 220		Thr Gln	Thr 225
55	Tyr Ile	Cys Asn	Val Asn 230	His Ly	s Pro Ser 235		: Lys Val	Asp 240
	Lys Lys	Val Glu	Pro Lys	Ser Cy	s Asp Lys	Thr His	: Thr	

PCT/US98/03337

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245	250	25
245	250	

(2) INFORMATION F	OR SEQ	ID	NO:67:
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- 5 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 159 amino acids
  - (B) TYPE: Amino Acid
  - (D) TOPOLOGY: Linear
- 10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:67:

Ser Gly Gly Gly Ser Gly Ser Gly Asp Phe Asp Tyr Glu Lys Met

15 Ala Asn Ala Asn Lys Gly Ala Met Thr Glu Asn Ala Asp Glu Asn 20 25 30

Ala Leu Gln Ser Asp Ala Lys Gly Lys Leu Asp Ser Val Ala Thr

20
Asp Tyr Gly Ala Ala Ile Asp Gly Phe Ile Gly Asp Val Ser Gly
50
55
60

Leu Ala Asn Gly Asn Gly Ala Thr Gly Asp Phe Ala Gly Ser Ser

5 70 75

Asn Ser Gln Met Ala Gln Val Gly Asp Gly Asp Asn Ser Pro Leu 80 85 90

Met Asn Asn Phe Arg Gln Tyr Leu Pro Ser Leu Pro Gln Ser Val 95 100 105

Glu Cys Arg Pro Phe Val Phe Ser Ala Gly Lys Pro Tyr Glu Phe

Ser Ile Asp Cys Asp Lys Ile Asn Leu Phe Arg Gly Val Phe Ala 125 130 135

Phe Leu Leu Tyr Val Ala Thr Phe Met Tyr Val Phe Ser Thr Phe
40 140 145 150

Ala Asn Ile Leu Arg Asn Lys Glu Ser 155 159

- 45 (2) INFORMATION FOR SEQ ID NO:68:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 780 base pairs
    - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
    - (D) TOPOLOGY: Linear
    - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:
- 55
  ATGAAAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT 50

WO<sup>-</sup>98/37200 PCT/US98/03337

	IGCIACAAC GCAIACGCIG AIAICCACAI GACCAGICO CONTOCIO DI													
_	TGTCCGCCTC TGTGGGCGAT AGGGTCACCA TCACCTGCAG GTCAAGTCAA													
5	AGCTTAGTAC ATGGTATAGG TAACACGTAT TTACACTGGT ATCAACAGAA 200													
	ACCAGGAAAA GCTCCGAAAC TACTGATTTA CAAAGTATCC AATCGATTCT 250													
10	CTGGAGTCCC TTCTCGCTTC TCTGGATCCG GTTCTGGGAC GGATTTCACT 300													
	CTGACCATCA GCAGTCTGCA GCCAGAAGAC TTCGCAACTT ATTACTGTTC 350													
15	ACAGAGTACT CATGTCCCGC TCACGTTTGG ACAGGGTACC AAGGTGGAGA 400													
13	TCAAACGAAC TGTGGCTGCA CCATCTGTCT TCATCTTCCC GCCATCTGAT 450													
	GAGCAGTTGA AATCTGGAAC TGCTTCTGTT GTGTGCCTGC TGAATAACTT 500													
20	CTATCCCAGA GAGGCCAAAG TACAGTGGAA GGTGGATAAC GCCCTCCAAT 550													
	CGGGTAACTC CCAGGAGAGT GTCACAGAGC AGGACAGCAA GGACAGCACC 600													
25	TACAGCCTCA GCAGCACCCT GACGCTGAGC AAAGCAGACT ACGAGAAACA 650													
	CAAAGTCTAC GCCTGCGAAG TCACCCATCA GGGCCTGAGC TCGCCCGTCA 700													
	CAAAGAGCTT CAACAGGGGA GAGTGTTAAG CTGATCCTCT ACGCCGGACG 750													
30	CATCGTGGCC CTAGTACGCA ACTAGTCGTA 780													
	(2) INFORMATION FOR SEQ ID NO:69:													
35	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 242 amino acids</li><li>(B) TYPE: Amino Acid</li><li>(D) TOPOLOGY: Linear</li></ul>													
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:													
40	Met Lys Lys Asn Ile Ala Phe Leu Leu Ala Ser Met Phe Val Phe 1 5 10													
45	Ser Ile Ala Thr Asn Ala Tyr Ala Asp Ile Gln Met Thr Gln Ser 20 25 30													
	Pro Ser Ser Leu Ser Ala Ser Val Gly Asp Arg Val Thr Ile Thr 35 40 45													
50	Cys Arg Ser Ser Gln Ser Leu Val His Gly Ile Gly Asn Thr Tyr 50 55 60													
	Leu His Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys Leu Leu 65 70 79													

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Ile Tyr Lys Val Ser Asn Arg Phe Ser Gly Val Pro Ser Arg Phe

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					80					85					90
	Ser G	iy	Ser	Gly	Ser 95	Gly	Thr	Asp	Phe	Thr 100	Leu	Thr	Ile	Ser	Ser 105
5	Leu G	ln	Pro	Glu	Asp 110	Phe	Ala	Thr	Tyr	Tyr 115	Cys	Ser	Gln	Ser	Thr 120
10	His V	/al	Pro	Leu	Thr 125	Phe	Gly	Gln	Gly	Thr 130	Lys	Val	Glu	Ile	Lys 135
	Arg T	Chr	Val	Ala	Ala 140	Pro	Ser	Val	Phe	Ile 145	Phe	Pro	Pro	Ser	Asp 150
15	Glu G	Sln	Leu	Lys	Ser 155	Gly	Thr	Ala	Ser	Val 160	Val	Cys	Leu	Leu	Asn 165
	Asn I	Phe	Tyr	Pro	Arg 170	Glu	Ala	Lys	Val	Gln 175	Trp	Lys	Val	Asp	Asn 180
20	Ala l	Leu	Gln	Ser	Gly 185	Asn	Ser	Gln	Glu	Ser 190	Val	Thr	Glu	Gln	Asp 195
25	Ser 1	Lys	Asp	Ser	Thr 200	Tyr	Ser	Leu	Ser	Ser 205	Thr	Leu	Thr	Leu	Ser 210
	Lys i	Ala	Asp	Tyr	Glu 215		His	Lys	Val	Туг 220		Cys	Glu	Val	Thr 225
30	His	Gln	Gly	Leu	Ser 230		Pro	Val	Thr	Lys 235		Phe	Asn	Arg	Gly 240
25	Glu	Cys 242													
35	(2) I	NFO	RMAT	ON	FOR	SEQ	ID N	10:70	:						
40	(i	(. (:	A) L B) T		H: 2 Ami	.no <i>F</i>									
	(xi	.) S	EQUE	NCE	DESC	RIPT	: NOI	SEC	O ID	NO: 7	70:				
45	Met 1	Lys	Lys	a Asr	ı Ile		a Phe	e Lev	ı Leı	ı Ala		. Met	: Phe	val	Phe 15
50	Ser	Ile	Ala	Thr	2 Ası		а Туі	c Ala	a Glv	ı Val		ı Let	ı Val	Glu	Ser 30
50	Gly	Gly	gly	/ Lev	ı Va:		n Pro	o Gly	y Gl	y Se:		ı Arç	g Lev	ı Seı	Cys 45
55	Ala	Ala	. Sei	c Gly	у Ту: 5	_	r Pho	e Se:	r Se	r Hi:		r Me	t His	s Trp	Val 60

Lys Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Gly Tyr Ile Asp

	•				65	_				70					75
5	Pro	Ser	Asn	Gly	Glu 80	Thr	Thr	Tyr	Asn	Gln 85	Lys	Phe	Lys	Gly	Arg 90
	Phe	Thr	Leu	Ser	Arg 95	Asp	Asn	Ser	Lys	Asn 100	Thr	Ala	Tyr	Leu	Gln 105
10	Met	Asn	Ser	Leu	Arg 110	Ala	Glu	Asp	Thr	Ala 115	Val	Tyr	Tyr	Cys	Ala 120
15	Arg	Gly	Asp	Tyr	Arg 125	Tyr	Asn	Gly	Asp	Trp 130	Phe	Phe	Asp	Val	Trp 135
13	Gly	Gln	Gly	Thr	Leu 140	Val	Thr	Val	Ser	Ser 145	Ala	Ser	Thr	Lys	Gly 150
20	Pro	Ser	Val	Phe	Pro 155	Leu	Ala	Pro	Ser	Ser 160	Lys	Ser	Thr	Ser	Gly 165
	Gly	Thr	Ala	Ala	Leu 170	Gly	Cys	Leu	Val	Lys 175	Asp	Tyr	Phe	Pro	Glu 180
25	Pro	Val	Thr	Val	Ser 185	Trp	Asn	Ser	Gly	Ala 190	Leu	Thr	Ser	Gly	Val 195
30	His	Thr	Phe	Pro	Ala 200	Val	Leu	Gln	Ser	Ser 205	Gly	Leu	Tyr	Ser	Leu 210
30	Ser	Ser	Val	Val	Thr 215	Val	Pro	Ser	Ser	Ser 220	Leu	Gly	Thr	Gln	Thr 225
35	Tyr	Ile	Cys	Asn	Val 230	Asn	His	Lys	Pro	Ser 235	Asn	Thr	Lys	Val	Asp 240
	Lys	Lys	Val	Glu	Pro 245	Lys	Ser	Cys	Asp	Lys 250	Thr	His	Thr 253		
40	(2)	INFO	RMAT	ION I	FOR :	SEQ	ID N	0:71	:						
	. (:	(.	A) L	NCE ( ENGT: YPE:	H: 2	42 a	mino		ds					_	
45				OPOL							•				
	(x	i) S	EQUE	NCE :	DESC	RIPT	ION:	SEQ	ID	NO : 7	1:				
50	Met 1	-	Lys	Asn	Ile 5		Phe	Leu	Leu	Ala 10		Met	Phe	Val	Phe 15
	Ser	Ile	Ala	Thr	Asn 20		Tyr	Ala	Asp	Ile 25		Met	Thr	Gln	Ser 30
55	Pro	Ser	Ser	Leu	Ser 35		Ser	Val	Gly	Asp 40		Val	Thr	Ile	Thr 45

	Суѕ	Arg	Ser	Ser	Gln 50	Ser	Leu	Val	His	Gly 55	Ile	Gly	Ala	Thr	Tyr 60
5	Leu	His	Trp	Tyr	Gln 65	Gln	Lys	Pro	Gly	Lys 70	Ala	Pro	Lys	Leu	Leu 75
	Ile	Tyr	Lys	Val	Ser 80	Asn	Arg	Phe	Ser	Gly 85	Val	Pro	Ser	Arg	Phe 90
10	Ser	Gly	Ser	Gly	Ser 95	Gly	Thr	Asp	Phe	Thr 100	Leu	Thr	Ile	Ser	Ser 105
15	Leu	Gln	Pro	Glu	Asp 110	Phe	Ala	Thr	Tyr	туг 115	Cys	Ser	Gln	Ser	Thr 120
	His	Vål	Pro	Leu	Thr 125	Phe	Gly	Gln	Gly	Thr 130	Lys	Val	Glu	Ile	Lys 135
20	Arg	Thr	Val	Ala	Ala		Ser	Val	Phe	Ile 145	Phe	Pro	Pro	Ser	Asp 150
	Glu	Gln	Leu	Lys	Ser 155		Thr	Ala	Ser	Val 160	Val	Cys	Leu	Leu	Asn 165
25	Asr	n Phe	туг	Pro	Arg		ı Ala	Lys	Val	Gln 175	Trp	Lys	Val	Asp	Asn 180
30	Ala	a Lei	ı Glr	ser	Gly 185		n Ser	Glr	Glu	Ser 190	Val	Thr	Glu	Gln	195
	Se	r Lys	s Asp	o. Sei	7hr 200		s Ser	Lev	ı Ser	Ser 205	Thr	Lev	Thr	Lev	210
35	Ly	s Ala	a Asp	э Туг	c Glu 21		s His	s Lys	s Val	220	Ala	суя	s Glu	ı Val	225
	Hi	s Gl	n Gly	y Lei	u Sei 230		r Pro	o Vai	l Thi	235	s Ser	Phe	e Ası	a Arg	Gly 240
40	Gl	u Cy 24													
45	(2)						ID I								
		(1)	(A) (B)	LENG TYPE	TH:	45 a ino	ERIS mino Acid	aci							
50	(	(xi)					near TION		Q ID	NO:	72:				
	c?	/s Pr 1	o Pr	ю Су	s Pr	o Al	a Pr	o G1	u Le	u Le 1	u Gl .0	y Gl	y Ar	g Me	t Lys 15
55	G]	ln Le	eu Ġl	lu As	sp Ly	rs Va	al Gl	.u G	u Le	eu Le	u Se	r Ly	s As	n Ty	r His

20 25 30

Leu Glu Asn Glu Val Ala Arg Leu Lys Lys Leu Val Gly Glu Arg
35 40 45

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- (2) INFORMATION FOR SEQ ID NO:73:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 780 base pairs
    - (B) TYPE: Nucleic Acid
    - (C) STRANDEDNESS: Single
    - (D) TOPOLOGY: Linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:

15

10

ATGAAAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT 50 TGCTACAAAC GCATACGCTG ATATCCAGAT GACCCAGTCC CCGAGCTCCC 100 20 AGCTTAGTAC ATGGTATAGG TGCTACGTAT TTACACTGGT ATCAACAGAA 200 ACCAGGAAAA GCTCCGAAAC TACTGATTTA CAAAGTATCC AATCGATTCT 250 25 CTGGAGTCCC TTCTCGCTTC TCTGGATCCG GTTCTGGGAC GGATTTCACT 300 CTGACCATCA GCAGTCTGCA GCCAGAAGAC TTCGCAACTT ATTACTGTTC 350 30 ACAGAGTACT CATGTCCCGC TCACGTTTGG ACAGGGTACC AAGGTGGAGA 400 TCAAACGAAC TGTGGCTGCA CCATCTGTCT TCATCTTCCC GCCATCTGAT 450 35 GAGCAGTTGA AATCTGGAAC TGCTTCTGTT GTGTGCCTGC TGAATAACTT 500 CTATCCCAGA GAGGCCAAAG TACAGTGGAA GGTGGATAAC GCCCTCCAAT 550 CGGGTAACTC CCAGGAGAGT GTCACAGAGC AGGACAGCAA GGACAGCACC 600 40 TACAGCCTCA GCAGCACCCT GACGCTGAGC AAAGCAGACT ACGAGAAACA 650

CAAAGTCTAC GCCTGCGAAG TCACCCATCA GGGCCTGAGC TCGCCCGTCA 700

CAAAGAGCTT CAACAGGGGA GAGTGTTAAG CTGATCCTCT ACGCCGGACG 750

(2) INFORMATION FOR SEQ ID NO:74:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 927 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single

CATCGTGGCC CTAGTACGCA ACTAGTCGTA 780

55 (D) TOPOLOGY: Linear

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:74:

"我们我们的"我们的"我们会看我"的变形。

	AAAAGGGTAT	CTAGAGGTTG	AGGTGATTTT	ATGAAAAAGA	ATATCGCATT	50
5	TCTTCTTGCA	TCTATGTTCG	TTTTTTCTAT	TGCTACAAAC	GCGTACGCTG	100
	AGGTTCAGCT	AGTGCAGTCT	GGCGGTGGCC	TGGTGCAGCC	AGGGGGCTCA	150
10	CTCCGTTTGT	CCTGTGCAGC	TTCTGGCTAC	TCCTTCTCGA	GTCACTATAT	200
	GCACTGGGTC	CGTCAGGCCC	CGGGTAAGGG	CCTGGAATGG	GTTGGATATA	250
	TTGATCCTTC	CAATGGTGAA	ACTACGTATA	ATCAAAAGTT	CAAGGGCCGT	300
15	TTCACTTTAT	CTCGCGACAA	СТССАААААС	ACAGCATACC	TGCAGATGAA	350
	CAGCCTGCGT	GCTGAGGACA	CTGCCGTCTA	TTACTGTGCA	AGAGGGGATT	400
20	ATCGCTACAA	TGGTGACTGG	TTCTTCGACG	TCTGGGGTCA	AGGAACCCTG	450
	GTCACCGTCT	CCTCGGCCTC	CACCAAGGGC	CCATCGGTCT	TCCCCCTGGC	500
	ACCCTCCTCC	AAGAGCACCT	CTGGGGGCAC	AGCGGCCCTG	GGCTGCCTGG	550
25	TCAAGGACTA	CTTCCCCGAA	CCGGTGACGG	TGTCGTGGAA	CTCAGGCGCC	600
	CTGACCAGCG	GCGTGCACAC	: CTTCCCGGCT	GTCCTACAGT	CCTCAGGACT	650
30	CTACTCCCTC	AGCAGCGTGG	TGACCGTGCC	CTCCAGCAGC	TTGGGCACCC	700
	AGACCTACAT	CTGCAACGT	AATCACAAGO	CCAGCAACAC	CAAGGTCGAC	750
25	AAGAAAGTT	G AGCCCAAAT	TTGTGACAA	A ACTCACACAT	GCCCGCCGTC	800
35	CCCAGCACC	A GAACTGCTG	GCGGCCGCA	r gaaacagct	A GAGGACAAGO	850
	TCGAAGAGC	r ACTCTCCAA	AACTACCAC	TAGAGAATG	A AGTGGCAAG	A 900
40	CTCAAAAAG	TTGTCGGGG	A GCGCTAA 9	27		

(2) INFORMATION FOR SEQ ID NO:75:

45

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 298 amino acids
  - (B) TYPE: Amino Acid
  - (D) TOPOLOGY: Linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:75:

Met Lys Lys Asn Ile Ala Phe Leu Leu Ala Ser Met Phe Val Phe
1 5 10 15

Ser Ile Ala Thr Asn Ala Tyr Ala Glu Val Gln Leu Val Gln Ser 55 20 25 30

	Gly	Gly	Gly	Leú	Val 35	Gln	Pro	Gly	Gly	Ser 40	Leu	Arg	Leu	Ser	Cys 45
5	Ala	Ala	Ser	Gly	Tyr 50	Ser	Phe	Ser	Ser	His 55	Tyr	Met	His	Trp	Val 60
	Arg	Gln	Ala	Pro	Gly 65	Lys	Gly	Leu	Glu	Trp 70	Val	Gly	Tyr	Ile	Asp 75
10	Pro	Ser	Asn	Gly	Glu 80	Thr	Thr	туr	Asn	Gln 85	Lys	Phe	Lys	Gly	Arg 90
16	Phe	Thr	Leu	Ser	Arg 95	Asp	Asn	Ser	Lys	Asn 100	Thr	Ala	Tyr	Leu	Gln 105
15	Met	Asn	Ser	Leu	Arg 110	Ala	Glu	Asp	Thr	Ala 115	Val	Tyr	Tyr	Cys	Ala 120
20	Arg	Gly	Asp	Tyr	Arg 125	Tyr	Asn	Gly	Asp	Trp 130	Phe	Phe	Asp	Val	Trp 135
	Gly	Gln	Gly	Thr	Leu 140	Val	Thr	Val	Ser	Ser 145	Ala	Ser	Thr	Lys	Gly 150
25	Pro	Ser	Val	Phe	Pro 155	Leu	Ala	Pro	Ser	Ser 160	Lys	Ser	Thr	Ser	Gly 165
20	Gly	Thr	Ala	Ala	Leu 170	Gly	Cys	Leu	Val	Lys 175	Asp	Tyr	Phe	Pro	Glu 180
30	Pro	Val	Thr	Val	Ser 185	Trp	Asn	Ser	Gly	Ala 190	Leu	Thr	Ser	Gly	Val 195
35	His	Thr	Phe	Pro	Ala 200	Val	Leu	Gln	Ser	Ser 205	Gly	Leu	Tyr	Ser	Leu 210
	Ser	Ser	Val	Val	Thr 215	Val	Pro	Ser	Ser	Ser 220		Gly	Thr	Gln	Thr 225
40	Tyr	Ile	Cys	Asn	Val 230	Asn	His	Lys	Pro	Ser 235		Thr	Lys	Val	Asp 240
45	Lys	Lys	Val	Glu	Pro 245		Ser	Cys	Asp	Lys 250		His	Thr	Cys	Pro 255
43	Pro	Cys	Pro	Ala	Pro 260		Leu	Leu	Gly	Gly 265		Met	Lys	Gln	Leu 270
50	Glu	Asp	Lys	Val	Glu 275		Leu	Leu	Ser	Lys 280		туг	His	Leu	Glu 285
	Asn	Glu	Val	Ala	Arg 290		Lys	Lys	. Leu	Val 295		Glu	Arg 298		
55	(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	10:76	i :						

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 6563 base pairs
  - (B) TYPE: Nucleic Acid
  - (C) STRANDEDNESS: Single
- 5 (D) TOPOLOGY: Linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:76:
- GAATTCAACT TCTCCATACT TTGGATAAGG AAATACAGAC ATGAAAAATC 50 10 TCATTGCTGA GTTGTTATTT AAGCTTGCCC AAAAAGAAGA AGAGTCGAAT 100 GAACTGTGTG CGCAGGTAGA AGCTTTGGAG ATTATCGTCA CTGCAATGCT 150 15 TCGCAATATG GCGCAAAATG ACCAACAGCG GTTGATTGAT CAGGTAGAGG 200 GGGCGCTGTA CGAGGTAAAG CCCGATGCCA GCATTCCTGA CGACGATACG 250 GAGCTGCTGC GCGATTACGT AAAGAAGTTA TTGAAGCATC CTCGTCAGTA 300 20 AAAAGTTAAT CTTTTCAACA GCTGTCATAA AGTTGTCACG GCCGAGACTT 350 ATAGTCGCTT TGTTTTTATT TTTTAATGTA TTTGTAACTA GAATTCGAGC 400 25 TCGGTACCCG GGGATCCTCT CGAGGTTGAG GTGATTTTAT GAAAAAGAAT 450 ATCGCATTTC TTCTTGCATC TATGTTCGTT TTTTCTATTG CTACAAACGC 500 ATACGCTGAT ATCCAGATGA CCCAGTCCCC GAGCTCCCTG TCCGCCTCTG 550 30 TGGGCGATAG GGTCACCATC ACCTGCAGGT CAAGTCAAAG CTTAGTACAT 600 GGTATAGGTG CTACGTATTT ACACTGGTAT CAACAGAAAC CAGGAAAAGC 650 35 TCCGAAACTA CTGATTTACA AAGTATCCAA TCGATTCTCT GGAGTCCCTT 700 CTCGCTTCTC TGGATCCGGT TCTGGGACGG ATTTCACTCT GACCATCAGC 750 AGTCTGCAGC CAGAAGACTT CGCAACTTAT TACTGTTCAC AGAGTACTCA 800 40 TGTCCCGCTC ACGTTTGGAC AGGGTACCAA GGTGGAGATC AAACGAACTG 850 TGGCTGCACC ATCTGTCTTC ATCTTCCCGC CATCTGATGA GCAGTTGAAA 900 45 TCTGGAACTG CTTCTGTTGT GTGCCTGCTG AATAACTTCT ATCCCAGAGA 950 GGCCAAAGTA CAGTGGAAGG TGGATAACGC CCTCCAATCG GGTAACTCCC 1000 AGGAGAGTGT CACAGAGCAG GACAGCAAGG ACAGCACCTA CAGCCTCAGC 1050 50 AGCACCCTGA CGCTGAGCAA AGCAGACTAC GAGAAACACA AAGTCTACGC 1100 CTGCGAAGTC ACCCATCAGG GCCTGAGCTC GCCCGTCACA AAGAGCTTCA 1150 55 ACAGGGGAGA GTGTTAAGCT GATCCTCTAC GCCGGACGCA TCGTGGCCCT 1200

	AGTACGCAAC	TAGTCGTAAA	AAGGGTATCT	AGAGGTTGAG	GTGATTTTAT	1250
_	GAAAAGAAT	ATCGCATTTC	TTCTTGCATC	TATGTTCGTT	TTTTCTATTG	1300
5	CTACAAACGC	GTACGCTGAG	GTTCAGCTAG	TGCAGTCTGG	CGGTGGCCTG	1350
	GTGCAGCCAG	GGGGCTCACT	CCGTTTGTCC	TGTGCAGCTT	CTGGCTACTC	1400
10	CTTCTCGAGT	CACTATATGC	ACTGGGTCCG	TCAGGCCCCG	GGTAAGGGCC	1450
	TGGAATGGGT	TGGATATATT	GATCCTTCCA	ATGGTGAAAC	TACGTATAAT	1500
	CAAAAGTTCA	AGGGCCGTTT	CACTTTATCT	CGCGACAACT	CCAAAAACAC	1550
15	AGCATACCTG	CAGATGAACA	GCCTGCGTGC	TGAGGACACT	GCCGTCTATT	1600
	ACTGTGCAAG	AGGGGATTAT	CGCTACAATG	GTGACTGGTT	CTTCGACGTC	1650
20	TGGGGTCAAG	GAACCCTGGT	CACCGTCTCC	TCGGCCTCCA	CCAAGGGCCC	1700
	ATCGGTCTTC	CCCCTGGCAC	CCTCCTCCAA	GAGCACCTCT	GGGGCACAG	1750
25	CGGCCCTGGG	CTGCCTGGTC	AAGGACTACT	TCCCCGAACC	GGTGACGGTG	1800
23	TCGTGGAACT	CAGGCGCCCT	GACCAGCGGC	GTGCACACCT	TCCCGGCTGT	1850
	CCTACAGTCC	TCAGGACTCT	ACTCCCTCAG	CAGCGTGGTG	ACCGTGCCCT	1900
30	CCAGCAGCTT	GGGCACCCAG	ACCTACATCT	GCAACGTGAA	TCACAAGCCC	1950
	AGCAACACCA	AGGTCGACAA	GAAAGTTGAG	CCCAAATCTT	GTGACAAAAC	2000
35	TCACACATGC	CCGCCGTGCC	CAGCACCAGA	ACTGCTGGGC	GGCCGCATGA	2050
<i>33</i>	AACAGCTAGA	GGACAAGGTC	GAAGAGCTAC	TCTCCAAGAA	CTACCACCTA	2100
	GAGAATGAAG	TGGCAAGACT	CAAAAAGCTT	GTCGGGGAGC	GCTAAGCATG	2150
40	CGACGGCCCT	AGAGTCCCTA	ACGCTCGGTT	GCCGCCGGGC	GTTTTTATT	2200
	GTTAACTCAT	GTTTGACAGC	TTATCATCGA	TAAGCTTTAA	TGCGGTAGTT	2250
45	TATCACAGTT	AAATTGCTAA	CGCAGTCAGG	CACCGTGTAT	GAAATCTAAC	2300
43	AATGCGCTCA	TCGTCATCCT	CGGCACCGTC	: ACCCTGGATG	CTGTAGGCAT	2350
	AGGCTTGGTT	ATGCCGGTAC	TGCCGGGCCT	CTTGCGGGAT	ATCGTCCATT	2400
50	CCGACAGCAT	CGCCAGTCAC	TATGGCGTGC	TGCTAGCGCT	ATATGCGTTG	2450
	ATGCAATTTC	TATGCGCACC	CGTTCTCGGA	A GCACTGTCCC	ACCGCTTTGG	2500
55	CCGCCGCCCA	GTCCTGCTCG	CTTCGCTACT	TGGAGCCACT	T ATCGACTACG	2550
JJ	ССАТСАТСС	. האכרארארר	GTCCTGTGG	TCCTCTACGO	CGGACGCATC	2600

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GTGGCCGGCA TCACCGGCGC CACAGGTGCG GTTGCTGGCG CCTATATCGC 2650 CGACATCACC GATGGGGAAG ATCGGGCTCG CCACTTCGGG CTCATGAGCG 2700 5 CTTGTTTCGG CGTGGGTATG GTGGCAGGCC CCGTGGCCGG GGGACTGTTG 2750 GGCGCCATCT CCTTGCACGC ACCATTCCTT GCGGCGGCGG TGCTCAACGG 2800 CCTCAACCTA CTACTGGGCT GCTTCCTAAT GCAGGAGTCG CATAAGGGAG 2850 10 AGCGTCGTCC GATGCCCTTG AGAGCCTTCA ACCCAGTCAG CTCCTTCCGG 2900 TGGGCGCGGG GCATGACTAT CGTCGCCGCA CTTATGACTG TCTTCTTTAT 2950 15 CATGCAACTC GTAGGACAGG TGCCGGCAGC GCTCTGGGTC ATTTTCGGCG 3000 AGGACCGCTT TCGCTGGAGC GCGACGATGA TCGGCCTGTC GCTTGCGGTA 3050 TTCGGAATCT TGCACGCCCT CGCTCAAGCC TTCGTCACTG GTCCCGCCAC 3100 20 CAAACGTTTC GGCGAGAAGC AGGCCATTAT CGCCGGCATG GCGGCCGACG 3150 CGCTGGGCTA CGTCTTGCTG GCGTTCGCGA CGCGAGGCTG GATGGCCTTC 3200 25 CCCATTATGA TTCTTCTCGC TTCCGGCGGC ATCGGGATGC CCGCGTTGCA 3250 GGCCATGCTG TCCAGGCAGG TAGATGACGA CCATCAGGGA CAGCTTCAAG 3300 GATCGCTCGC GGCTCTTACC AGCCTAACTT CGATCACTGG ACCGCTGATC 3350 30 GTCACGGCGA TTTATGCCGC CTCGGCGAGC ACATGGAACG GGTTGGCATG 3400 GATTGTAGGC GCCGCCCTAT ACCTTGTCTG CCTCCCCGCG TTGCGTCGCG 3450 35 GTGCATGGAG CCGGGCCACC TCGACCTGAA TGGAAGCCGG CGGCACCTCG 3500 CTAACGGATT CACCACTCCA AGAATTGGAG CCAATCAATT CTTGCGGAGA 3550 ACTGTGAATG CGCAAACCAA CCCTTGGCAG AACATATCCA TCGCGTCCGC 3600 40 CATCTCCAGC AGCCGCACGC GGCGCATCTC GGGCAGCGTT GGGTCCTGGC 3650 CACGGGTGCG CATGATCGTG CTCCTGTCGT TGAGGACCCG GCTAGGCTGG 3700 45 CGGGGTTGCC TTACTGGTTA GCAGAATGAA TCACCGATAC GCGAGCGAAC 3750 GTGAAGCGAC TGCTGCTGCA AAACGTCTGC GACCTGAGCA ACAACATGAA 3800 TGGTCTTCGG TTTCCGTGTT TCGTAAAGTC TGGAAACGCG GAAGTCAGCG 3850 50 CCCTGCACCA TTATGTTCCG GATCTGCATC GCAGGATGCT GCTGGCTACC 3900 CTGTGGAACA CCTACATCTG TATTAACGAA GCGCTGGCAT TGACCCTGAG 3950 55 TGATTTTCT CTGGTCCCGC CGCATCCATA CCGCCAGTTG TTTACCCTCA 4000

	CAACGTTCCA	GTAACCGGGC	ATGTTCATCA	TCAGTAACCC	GTATCGTGAG	4050
	CATCCTCTCT	CGTTTCATCG	GTATCATTAC	CCCCATGAAC	AGAAATTCCC	4100
5	CCTTACACGG	AGGCATCAAG	TGACCAAACA	GGAAAAAACC	GCCCTTAACA	4150
	TGGCCCGCTT	TATCAGAAGC	CAGACATTAA	CGCTTCTGGA	GAAACTCAAC	4200
10	GAGCTGGACG	CGGATGAACA	GGCAGACATC	TGTGAATCGC	TTCACGACCA	4250
	CGCTGATGAG	CTTTACCGCA	GCTGCCTCGC	GCGTTTCGGT	GATGACGGTG	4300
1.6	AAAACCTCTG	ACACATGCAG	CTCCCGGAGA	CGGTCACAGC	TTGTCTGTAA	4350
15	GCGGATGCCG	GGAGCAGACA	AGCCCGTCAG	GGCGCGTCAG	CGGGTGTTGG	4400
	CGGGTGTCGG	GGCGCAGCCA	TGACCCAGTC	ACGTAGCGAT	AGCGGAGTGT	4450
20	ATACTGGCTT	AACTATGCGG	CATCAGAGCA	GATTGTACTG	AGAGTGCACC	4500
	ATATGCGGTG	TGAAATACCG	CACAGATGCG	TAAGGAGAAA	ATACCGCATC	4550
25	AGGCGCTCTT	CCGCTTCCTC	GCTCACTGAC	TCGCTGCGCT	CGGTCGTTCG	4600
23	GCTGCGGCGA	GCGGTATCAG	CTCACTCAAA	GGCGGTAATA	CGGTTATCCA	4650
•	CAGAATCAGG	GGATAACGCA	GGAAAGAACA	TGTGAGCAAA	AGGCCAGCAA	4700
30	AAGGCCAGGA	ACCGTAAAAA	GGCCGCGTTG	CTGGCGTTTT	TCCATAGGCT	4750
	CCGCCCCCT	GACGAGCATC	ACAAAAATCG	ACGCTCAAGT	CAGAGGTGGC	4800
3 <i>5</i>	GAAACCCGAC	AGGACTATAA	AGATACCAGG	CGTTTCCCCC	TGGAAGCTCC	4850
33	CTCGTGCGCT	CTCCTGTTCC	GACCCTGCCG	CTTACCGGAT	ACCTGTCCGC	4900
	CTTTCTCCCT	TCGGGAAGCG	TGGCGCTTTC	TCATAGCTCA	CGCTGTAGGT	4950
40	ATCTCAGTTC	GGTGTAGGTC	GTTCGCTCCA	AGCTGGGCTG	TGTGCACGAA	5000
	CCCCCGTTC	AGCCCGACCG	CTGCGCCTTA	TCCGGTAACT	ATCGTCTTGA	5050
45	GTCCAACCCG	GTAAGACACG	ACTTATCGCC	ACTGGCAGCA	GCCACTGGTA	5100
	ACAGGATTAG	CAGAGCGAGG	TATGTAGGCG	GTGCTACAGA	GTTCTTGAAG	5150
	TGGTGGCCTA	ACTACGGCTA	CACTAGAAGG	ACAGTATTTG	GTATCTGCGC	5200
50	TCTGCTGAAG	CCAGTTACCT	TCGGAAAAAG	AGTTGGTAGC	TCTTGATCCG	5250
	GCAAACAAAC	CACCGCTGGT	AGCGGTGGTT	TTTTTGTTTG	CAAGCAGCAG	5300
55	ATTACGCGCA	GAAAAAAAGG	ATCTCAAGAA	GATCCTTTGA	. TCTTTTCTAC	5350
23	GGGGTCTGAC	GCTCAGTGGA	ACGAAAACTC	ACGTTAAGGG	ATTTTGGTCA	5400

TGAGATTATC AAAAAGGATC TTCACCTAGA TCCTTTTAAA TTAAAAATGA 5450 AGTTTTAAAT CAATCTAAAG TATATATGAG TAAACTTGGT CTGACAGTTA 5500 5 CCAATGCTTA ATCAGTGAGG CACCTATCTC AGCGATCTGT CTATTTCGTT 5550 CATCCATAGT TGCCTGACTC CCCGTCGTGT AGATAACTAC GATACGGGAG 5600 GGCTTACCAT CTGGCCCCAG TGCTGCAATG ATACCGCGAG ACCCACGCTC 5650 10 ACCGGCTCCA GATTTATCAG CAATAAACCA GCCAGCCGGA AGGGCCGAGC 5700 GCAGAAGTGG TCCTGCAACT TTATCCGCCT CCATCCAGTC TATTAATTGT 5750 15 TGCCGGGAAG CTAGAGTAAG TAGTTCGCCA GTTAATAGTT TGCGCAACGT 5800 TGTTGCCATT GCTGCAGGCA TCGTGGTGTC ACGCTCGTCG TTTGGTATGG 5850 CTTCATTCAG CTCCGGTTCC CAACGATCAA GGCGAGTTAC ATGATCCCCC 5900 20 ATGTTGTGCA AAAAAGCGGT TAGCTCCTTC GGTCCTCCGA TCGTTGTCAG 5950 AAGTAAGTTG GCCGCAGTGT TATCACTCAT GGTTATGGCA GCACTGCATA 6000 25 ATTCTCTTAC TGTCATGCCA TCCGTAAGAT GCTTTTCTGT GACTGGTGAG 6050 TACTCAACCA AGTCATTCTG AGAATAGTGT ATGCGGCGAC CGAGTTGCTC 6100 TTGCCCGGCG TCAACACGGG ATAATACCGC GCCACATAGC AGAACTTTAA 6150 30 AAGTGCTCAT CATTGGAAAA CGTTCTTCGG GGCGAAAACT CTCAAGGATC 6200 TTACCGCTGT TGAGATCCAG TTCGATGTAA CCCACTCGTG CACCCAACTG 6250 35 ATCTTCAGCA TCTTTTACTT TCACCAGCGT TTCTGGGTGA GCAAAAACAG 6300 GAAGGCAAAA TGCCGCAAAA AAGGGAATAA GGGCGACACG GAAATGTTGA 6350 ATACTCATAC TCTTCCTTTT TCAATATTAT TGAAGCATTT ATCAGGGTTA 6400 40 TTGTCTCATG AGCGGATACA TATTTGAATG TATTTAGAAA AATAAACAAA 6450 TAGGGGTTCC GCGCACATTT CCCCGAAAAG TGCCACCTGA CGTCTAAGAA 6500 45 ACCATTATTA TCATGACATT AACCTATAAA AATAGGCGTA TCACGAGGCC 6550 CTTTCGTCTT CAA 6563

#### WE CLAIM:

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A conjugate consisting essentially of one or more antibody fragments covalently attached to one or more nonproteinaceous polymer molecules, wherein the apparent size of the conjugate is at least about 500 kD.

- 2. The conjugate of claim 1, wherein the apparent size of the conjugate is at least about 800 kD.
- The conjugate of claim 1, wherein the apparent size of the conjugate is at least about 1,400 kD.
  - 4. The conjugate of claim 1, wherein the apparent size of the conjugate is at least about 1,800 kD.
    - 5. The conjugate of claim 1, wherein the apparent size of the conjugate is at least about 8 fold greater than the apparent size of the antibody fragment.
- 6. The conjugate of claim 5, wherein the apparent size of the conjugate is at least about 15 fold greater than the apparent size of the antibody fragment.
  - 7. The conjugate of claim 6, wherein the apparent size of the conjugate is at least about 25 fold greater than the apparent size of the antibody fragment.
- 25 8. The conjugate of claim 1, wherein the conjugate contains no more than one antibody fragment, and wherein the antibody fragment is selected from the group consisting of Fab, Fab', Fab'-SH, Fv, scFv and F(ab')<sub>2</sub>.
  - 9. The conjugate of claim 8 wherein the antibody fragment is F(ab')<sub>2</sub>.
  - 10. The conjugate of claim 1 wherein the antibody fragment is covalently attached to no more than about 10 nonproteinaceous polymer molecules.
- The conjugate of claim 10 wherein the antibody fragment is covalently attached to no more than about 5 nonproteinaceous polymer molecules.

12. The conjugate of claim 11 wherein the antibody fragment is covalently attached to no more than about 2 nonproteinaceous polymer molecules.

- 13. The conjugate of claim 12 wherein the antibody fragment is attached to no more than 1 nonproteinaceous polymer molecule.
  - 14. The conjugate of claim 12, wherein the antibody fragment comprises a heavy chain and a light chain derived from a parental antibody, wherein in the parental antibody the heavy and light chains are covalently linked by a disulfide bond between a cysteine residue in the light chain and a cysteine residue in the heavy chain, wherein in the antibody fragment the cysteine residue in the light or heavy chain is substituted with another amino acid and the cysteine residue in the opposite chain is covalently linked to a nonproteinaceous polymer molecule.
- 15. The conjugate of claim 8 wherein the antibody fragment is selected from the group consisting of Fab, Fab' and Fab'-SH.
  - 16. The conjugate of claim 15 wherein the antibody fragment is covalently attached to no more than I nonproteinaceous polymer molecule.
- 20 17. The conjugate of claim 16 wherein the nonproteinaceous polymer molecule in the conjugate is covalently attached to the hinge region of the antibody fragment.
  - 18. The conjugate of claim 1 wherein the nonproteinaceous polymer is a polyethylene glycol (PEG).
  - 19. The conjugate of claim 18 wherein the PEG has an average molecular weight of at least about 20 kD.
- The conjugate of claim 19 wherein the PEG has an average molecular weight of at least about 40 kD.
  - 21. The conjugate of claim 20 wherein the PEG is a single chain molecule.
  - 22. The conjugate of claim 20 wherein the PEG is a branched chain molecule.

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23. The conjugate of claim 19, wherein the conjugate contains no more than one antibody fragment, and wherein the antibody fragment is a F(ab')<sub>2</sub> and is covalently attached to no more than about 2 PEG molecules.

24. The conjugate of claim 19, wherein the conjugate contains no more than one antibody fragment, and wherein the antibody fragment is selected from the group consisting of Fab, Fab' and Fab'-SH and is covalently attached to no more than one PEG molecule.

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- The conjugate of claim 24 wherein the PEG molecule is covalently attached to the hinge region of the antibody fragment.
  - 26. The conjugate of claim 1 wherein the antibody fragment has an antigen binding site that binds to human IL-8.
- 15 27. The conjugate of claim 26, wherein the conjugate contains no more than one antibody fragment, wherein the antibody fragment is selected from the group consisting of Fab, Fab' and Fab'-SH, wherein the antibody fragment is covalently attached to no more than one nonproteinaceous polymer molecule, and wherein the nonproteinaceous polymer molecule is a polyethylene glycol having an actual molecular weight of at least about 30 kD.

28. The conjugate of claim 1 wherein the antibody fragment is humanized.

- 29. The conjugate of claim 1 wherein the conjugate contains no more than one antibody fragment.
  - 30. A composition comprising the conjugate of claim 1 and a carrier.
  - 31. The composition of claim 30 that is sterile.
- 30 32. A conjugate formed by one or more antibody fragments covalently attached to one or more nonproteinaceous polymer molecules, wherein the apparent size of the conjugate is at least about 500 kD, and wherein the molecular structure of the conjugate is free of other matter.
- 33. A conjugate formed by one or more antibody fragments covalently attached to one or more nonproteinaceous polymer molecules, wherein the apparent size of the conjugate is at least about 500 kD, wherein the antibody fragment incorporates a nonproteinaceous label free of any polymer, and wherein the molecular structure of the conjugate is free of other matter.

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34. The conjugate of claim 33 wherein the nonproteinaceous label is a radiolabel.

35. A polypeptide selected from the group consisting of: (1) a polypeptide that is an anti-IL-8 monoclonal antibody or antibody fragment comprising a light chain amino acid sequence comprising the complementarity determining regions of the light chain polypeptide amino acid sequence of Fig. 36; and (2) a polypeptide that is an anti-IL-8 monoclonal antibody or antibody fragment comprising a light chain amino acid sequence comprising the complementarity determining regions of the light chain polypeptide amino acid sequence of Fig. 45.

The polypeptide of claim 35, wherein the light chain amino acid sequence comprises the complementarity determining regions of the light chain polypeptide amino acid sequence of Fig. 45.

- The polypeptide of claim 35 that further comprises a heavy chain amino acid sequence comprising the complementarity determining regions of the heavy chain polypeptide amino acid sequence of Figs. 37A-37B.
  - 38. The polypeptide of claim 35 wherein the light chain amino acid sequence is selected from the group consisting of: (1) a light chain amino acid sequence comprising amino acids 1-219 of the light chain polypeptide amino acid sequence of Fig. 36; and (2) a light chain amino acid sequence comprising amino acids 1-219 of the light chain polypeptide amino acid sequence of Fig. 45.
  - 39. The polypeptide of claim 38 wherein the light chain amino acid sequence comprises amino acids 1-219 of the light chain amino acid sequence of Fig. 45.
  - 40. The polypeptide of claim 38 that further comprises a heavy chain amino acid sequence comprising amino acids 1-230 of the heavy chain polypeptide amino acid sequence of Figs. 37A-37B.
- The polypeptide of claim 40, wherein the heavy chain amino acid sequence is fused at its

  C-terminus to a leucine zipper amino acid sequence.
  - 42. The polypeptide of claim 41, wherein the leucine zipper sequence comprises amino acids 231-275 of the heavy chain polypeptide amino acid sequence of Figs. 37A-37B.
- 35 43. The polypeptide of claim 35 that is an antibody fragment selected from the group consisting of Fab, Fab', Fab'-SH, Fv, scFv and F(ab') 2.

44. The polypeptide of claim 38 that is a F(ab') 2 antibody fragment, wherein the antibody fragment comprises a first heavy chain amino acid sequence and a second heavy chain amino acid sequence each comprising amino acids 1-238 of the heavy chain polypeptide amino acid sequence of Figs. 37A-37B, and wherein each of the Cys residues at positions 231 and 234 in the first heavy chain amino acid sequence is in a disulfide linkage with the identical Cys residue in the second heavy chain amino acid sequence.

- 45. The polypeptide of claim 38 that is a Fab' or Fab'-SH antibody fragment, wherein the antibody fragment comprises a heavy chain amino acid sequence comprising amino acids 1-233 of the heavy chain polypeptide amino acid sequence of Fig. 53.
  - 46. The polypeptide of claim 35 that is an antibody.

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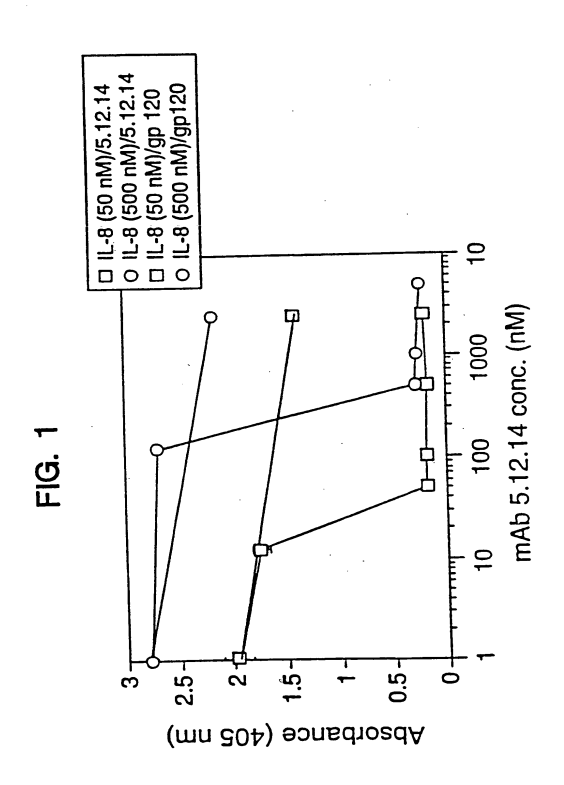
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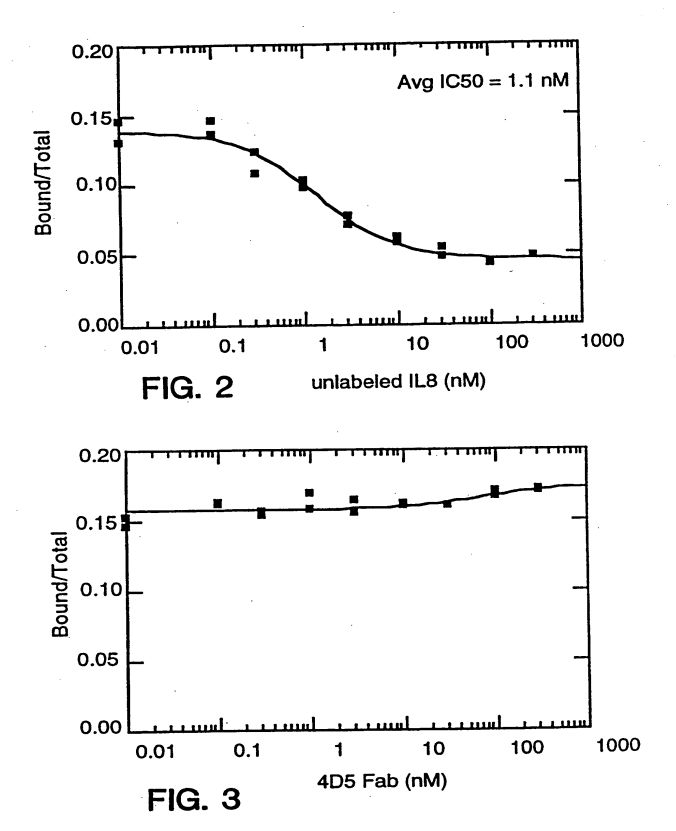
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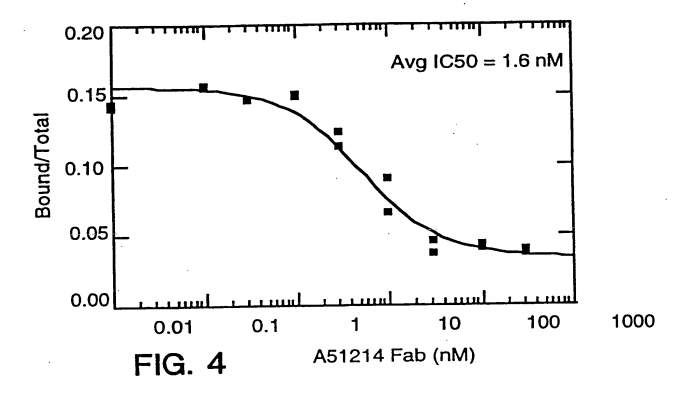
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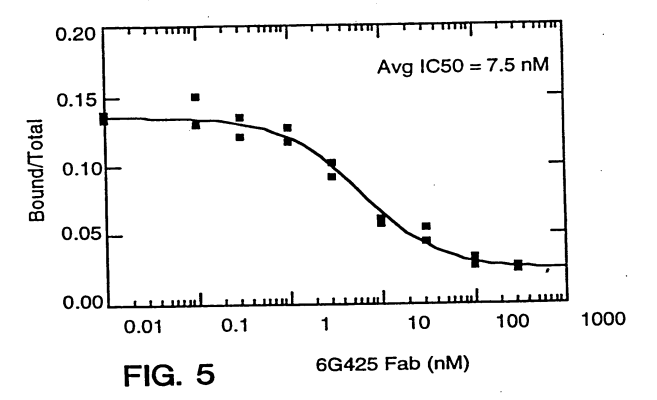
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- 47. A nucleic acid molecule that comprises a nucleic acid sequence encoding the polypeptide of claim 35.
- 48. An expression vector comprising the nucleic acid molecule of claim 47 operably linked to control sequences recognized by a host cell transfected with the vector.
  - 49. A host cell comprising the vector of claim 48.
- 50. A method of producing a polypeptide, comprising culturing the host cell of claim 49 under conditions wherein the nucleic acid sequence is expressed, thereby producing the polypeptide, and recovering the polypeptide from the host cell.
- 51. A composition comprising the polypeptide of claim 35 and a carrier.
  - 52. The composition of claim 51 that is sterile.









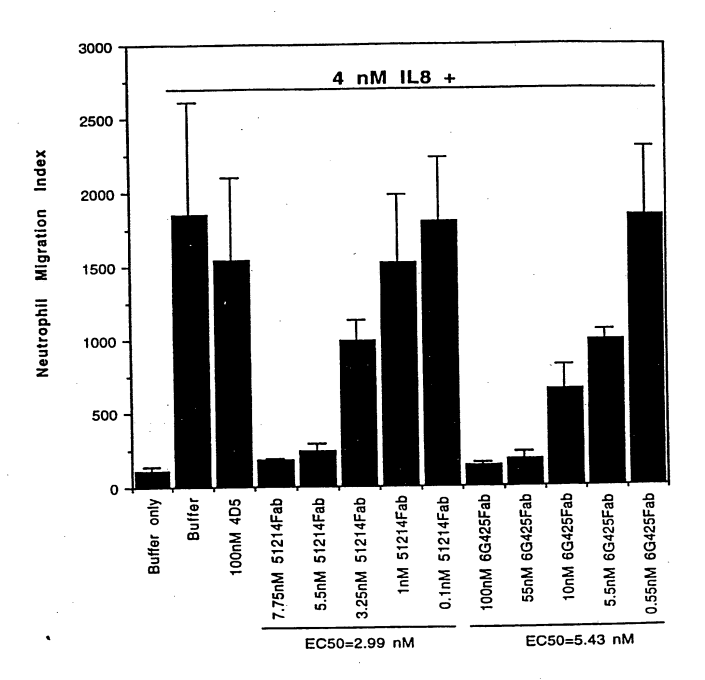


FIG. 6

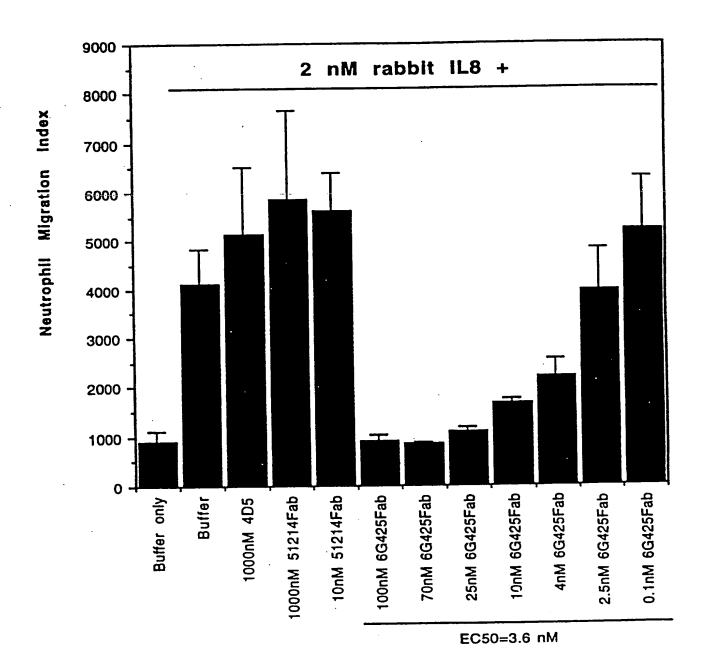
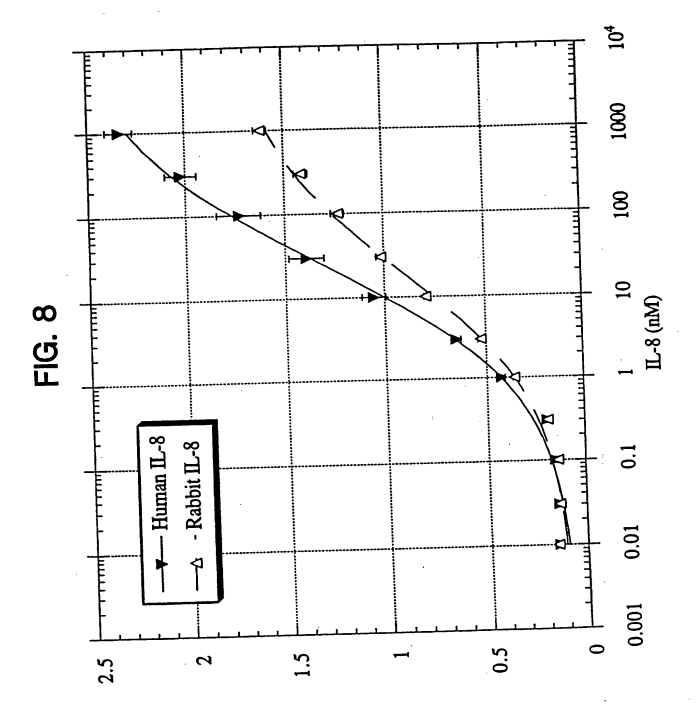
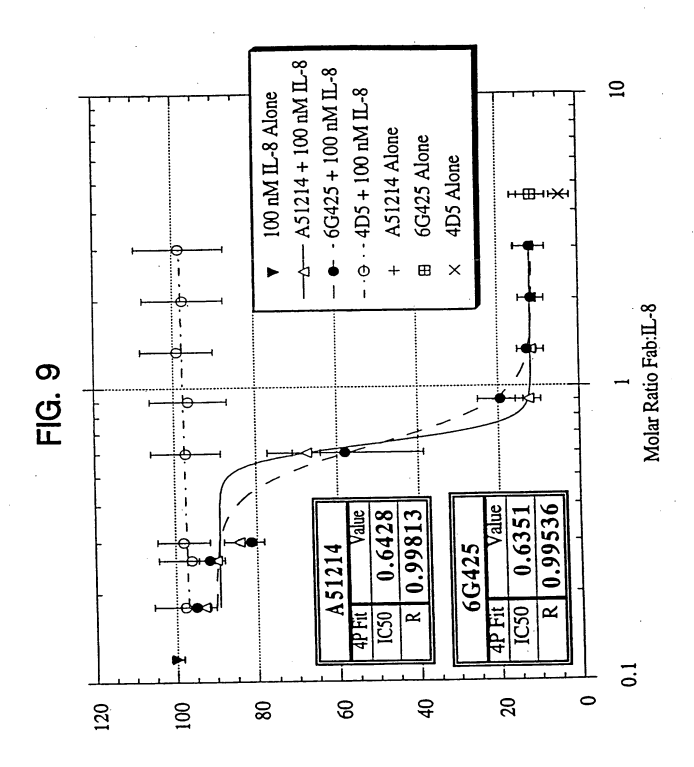


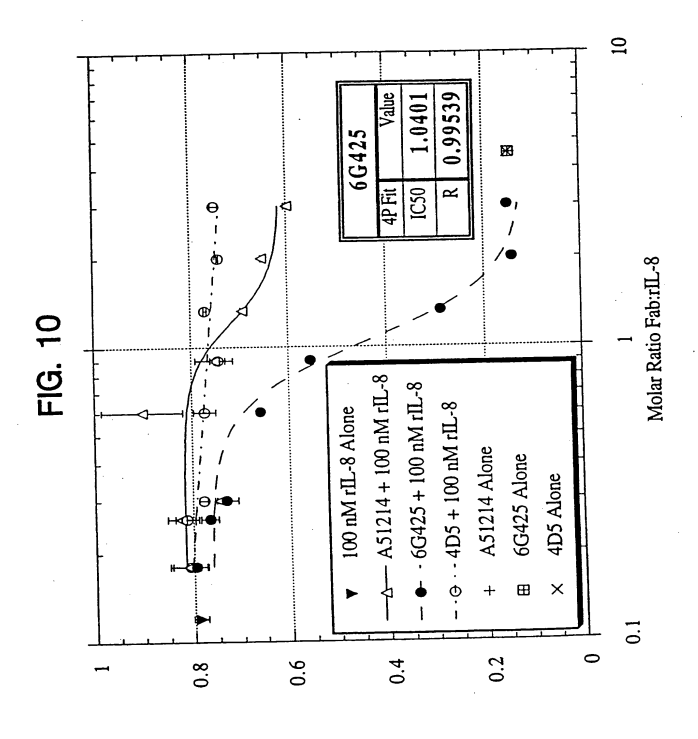
FIG. 7



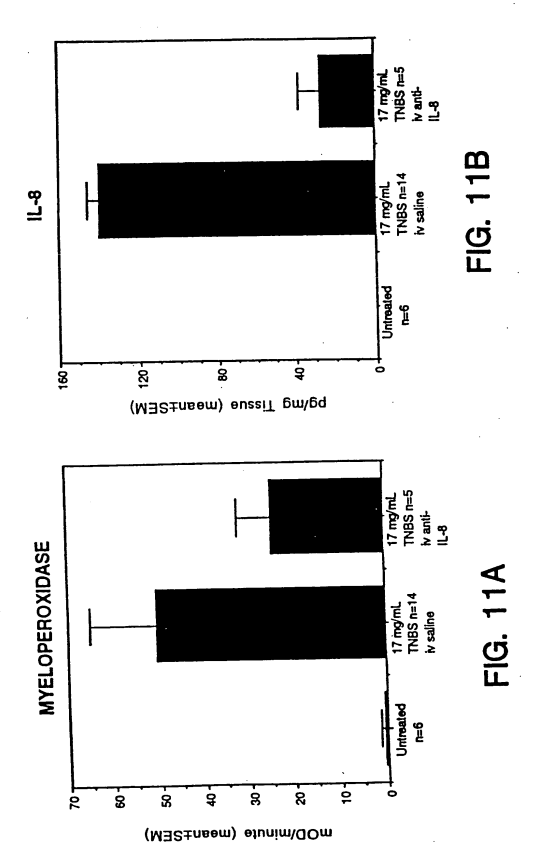
Absorbance (405 nm)



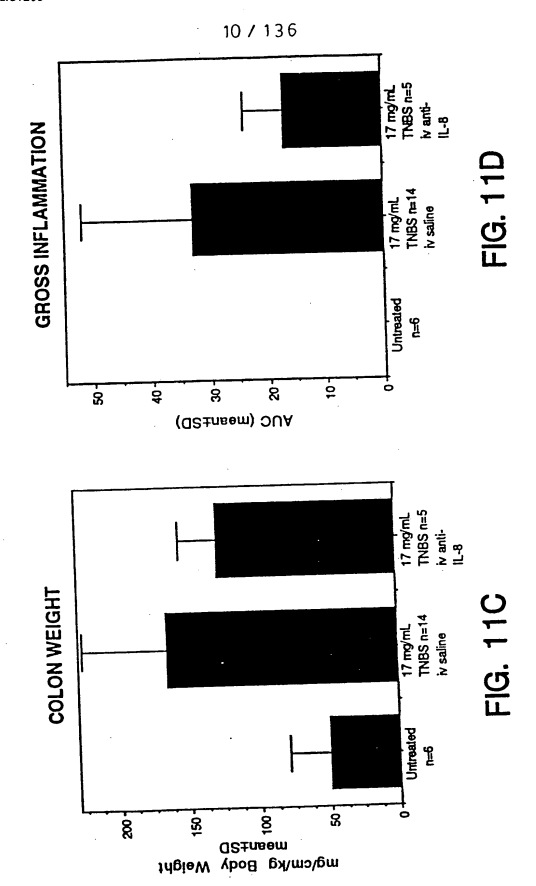
% IL-8-Stimulated Elastase Release



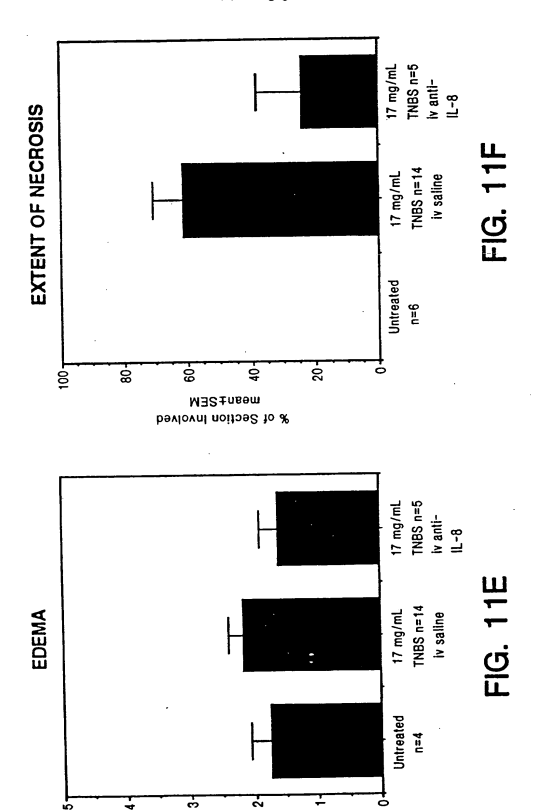
Absorbance (405 nm)



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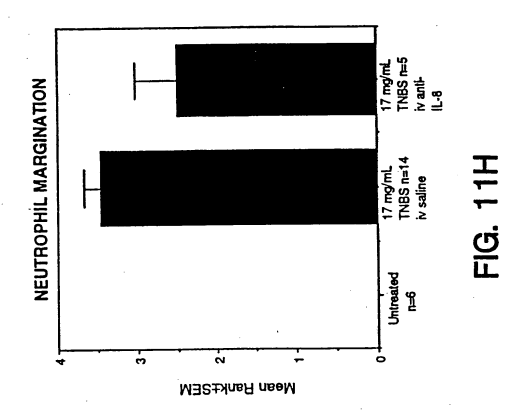


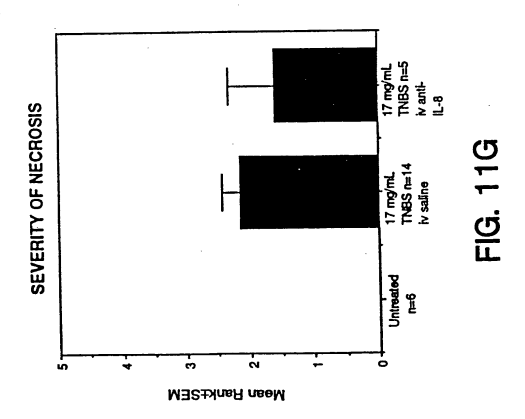
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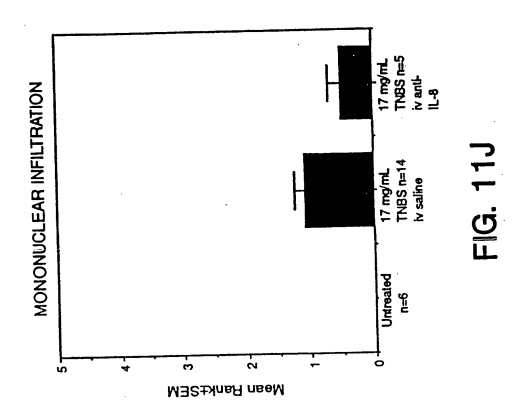
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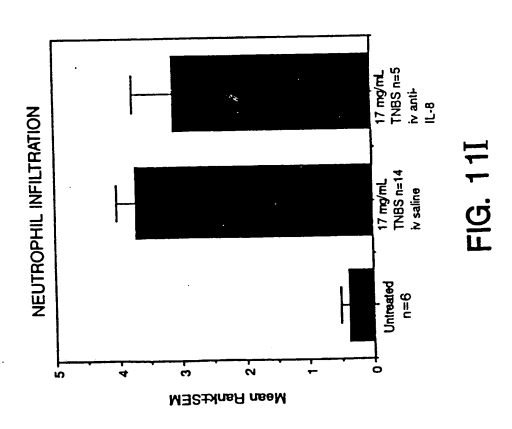
Mean Rank±SEM



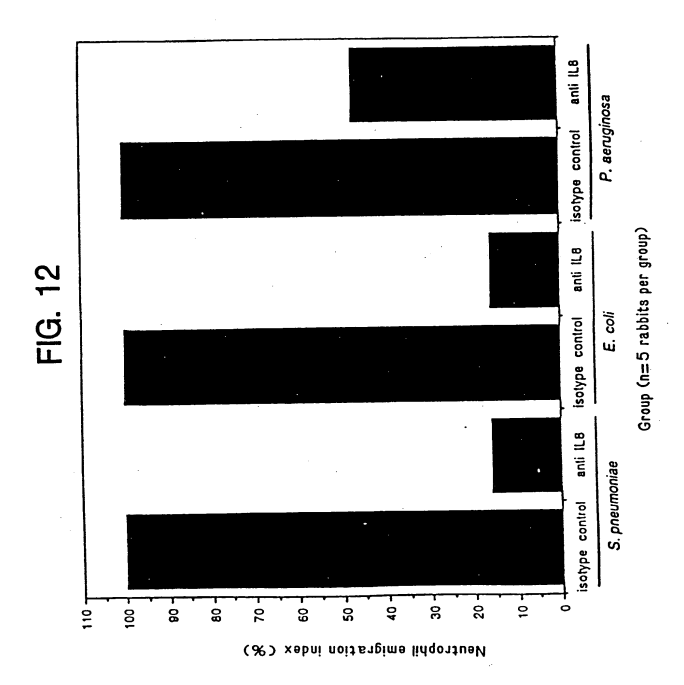


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Light Chain Primers:	_
MKLC-1, 22mer FIG. 1	] 3
5 · CAGTCCAACTGTTCAGGACGCC	: 3 <u>'</u>
MKLC-2, 22mer	
5' GTGCTGCTCATGCTGTAGGTGC	3'
MKLC-3, 23mer	
5' GAAGTTGATGTCTTGTGAGTGC	GC 3'
Heavy Chain Primers:	
IGG2AC-1, 24mer	
5' GCATCCTAGAGTCACCGAGGA	GCC 3
IGG2AC-2, 22mer	
5' CACTGGCTCAGGGAAATAACC	C 3'
IGG2AC-3, 22mer	
5' GGAGAGCTGGGAAGGTGTGCA	C 3'

5 '

# FIG. 14

Light chain forward primer

SL001A-2 35 mer

5 · ACAAACGCGTACGCT GACATCGTCATGACCCAGTC 3 · T T T A

Light chain reverse primer

SL001B 37 mer

5' GCTCTTCGAATG GTGGGAAGATGGATACAGTTGGTGC 3'

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Heavy chain forward primer

FIG. 15

SL002B 39 mer

5' CGATGGGCCCGG ATAGACCGATGGGGCTGTTGTTTTGGC 3'

T

G

Α

Heavy chain reverse primer

SL002B 39-MER

5' CGATGGGCCCGG ATAGACCGATGGGGCTGTTGTTTTGGC 3'

T

Α

G

GTCCCAGTCG CAGGGTCAGC > GACATTGTCA TGACACAGTC TCAAAAATTC ATGTCCACAT CAGTAGGAGA GTCATCCTCT Ö > TACAGGTGTA ß AGTTTTTAAG Ø CTGTAACAGT ACTGTGTCAG

ACAGAAACCA TGTCTTTGGT Ø CCTGGTATCA GGACCATAGT O 3 GAATGTGGGT ACTAATGTAG TGATTACATC CTTACACCCA GTCACCTGCA AGGCCAGTCA CAGTGGACGT TCCGGTCAGT ď 61

CDR #1

GATITACICG ICAICCIACC GGIACAGIGG AGICCCIGAI TCAGGGACTA AGTAGGATGG CCATGTCACC αļ \* CTAAATGAGC GATTTCGTGA CTAAAGCACT æ GGGCAATCTC CCCGTTAGAG Q

CDR #2

ACACGTCAGA TGTGCAGTCT > CCATCAGCCA GGTAGTCGGT TGGGACAGAT TTCACTCTCA AAGTGAGAGT ч E ACCCTGTCTA E G GCAGTGGATC CGTCACCTAG O Ŋ CGCTTCACAG GCGAAGTGTC 181 61

CAAGCCAGGA GTTCGGTCCT ტ TATAACATCT ATCCTCTCAC TAGGAGAGTG Ωı ATATTGTAGA CTGTCAGCAA GACAGTCGTT Ø GAAGACTTGG CAGACTATTT GTCTGATAAA CTTCTGAACC 241 81

CDR #3

CATCTTCCCA GTAGAAGGGT GCTGCACCAC CAACTGTATC CGACGTGGTG GTTGACATAG > Q, ACGGCCTGAT TGCCCGACTA æ GGGACCAAGC TGGAGTTGAA ACCTCAACTT CCCTGGTTCG 301 101

BstBI

361 CCATTCGAA GGTAAGCTT

回回

Д

FIG. 16

1	TTCTATTGCT	ACAAACGCGT	ACGCTGAGGT	GCAGCTGGTG	GAGTCTGGGG	GAGGCTTAGT
	<b>AAGATAACGA</b>	TGTTTGCGCA	TGCGACTCCA		CTCAGACCCC	CTCCGAATCA
1			E V	O r A	E S G G	G L V
61	CCCCCCTCCA	GGGTCCCTGA	AACTCTCCTG	TGCAGCCTCT	GGATTCATAT	TCAGTAGTTA
01	CCCCCACCT	CCCAGGGACT	TTGAGAGGAC	ACGTCGGAGA	CCTAAGTATA	AGTCATCAAT
12	P P G	G S L K			GFIF	SSY
13	PPG	א ע כ ט	H 5 C	A A D	34	* *
					CDR :	<b>#1</b>
121	TGGCATGTCT	TGGGTTCGCC	AGACTCCAGG	CAAGAGCCTG	GAGTTGGTCG	СААССАТТАА
	ACCGTACAGA	ACCCAAGCGG	TCTGAGGTCC	GTTCTCGGAC	CTCAACCAGC	GTTGGTAATT
33	G M S	WVRQ		K S L	E L V A	TIN
	* * *					* * *
181	TAATAATGGT	GATAGCACCT	ATTATCCAGA	CAGTGTGAAG	GGCCGATTCA	CCATCTCCCG
	ATTATTACCA	CTATCGTGGA	TAATAGGTCT		CCGGCTAAGT	
53	N N G	<u>D</u> S T Y	Y P D	s v K	G R F T	I S R
	<b>1</b> 2	* * * *	* * *	* * *		
		CDR #3	2			
241	AGACAATGCC	AAGAACACCC	TGTACCTGCA	AATGAGCAGT	CTGAAGTCTG	AGGACACAGC
	TCTGTTACGG	TTCTTGTGGG	ACATGGACGT			TCCTGTGTCG
73	D N A	K N T L	Y L Q	M S S	LKSE	D T A
301	CATGTTTTAC	TGTGCAAGAG	CCCTCATTAG	TTCGGCTACT	TGGTTTGGTT	ACTGGGGCCA
	GTACAAAATG	ACACGTTCTC				TGACCCCGGT
93	M F Y	C A R A		<u>sat</u>	W F G Y	, W G Q
		☆	<b>\$ \$</b>	<b>* * *</b>	* * * *	
			C	DR #3		
361	AGGGACTCTG	GTCACTGTCT	CTGCAGCCAA	AACAACAGCC	CCATCTGTCT GGTAGACAGA	
112		V T V S		T T A	P S V Y	
113	G T L	v i v s	AAA			
4 = =	ApaI					
411				r		,
400	TAGGCCC			ļ	FIG. 17	
130	P					

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# FIG. 18

VL.front	31-MER		
5' ACAA <u>ACGCGI</u> VL.rear 31-M	PACGCT <u>GATATC</u> GTCATGACAG ER	3'	
5' GCAGCATCAG	GCTC <u>TTCGAA</u> GCTCCAGCTTGG	3 '	
VH.front.SPE	21-MER		-
5' CCACTAGTA	CGCAAGTTCACG	. 3 '	
VH.rear 33-M	ER		
5 ' CATCCCCCC	TTGGTGGAGGCTGCAGAGACAG	rg	3

216

1	ATGAA(	GAAG:	A A	ATATO PATAO	:GCI	TTF FAA	TČTI AGAJ	<b>AGA</b>	CGT	AGA	TAC	CAAC	SC	AAAA	AAG	YI'A	ACGA	MGT	"I"IG
-23	M K	K 1	N	I	A	F	L	L	A	S	M	F	V	F	S	I	A	T	N
61	GCGTA	CGCT	G 1	TATA ATA1	CGT	CAT GTA	GACA CTGT	CAC	TCT CAGA	GTI	TTT	DAAT	CA '	TGTC ACAG	CAC GTG'	rag	AGTA TCA1	GGA CCI	GAC CTG
-3	A Y	A		I	V	M	T	Q	S	Q	K	F	M	S	T	S	V	G	D
121	AGGGT(	CAGC(	G 7	rcac( agrg(	CTG(	CAA GTT	GGC	CAGT	rcag agtc	AAT TTP	GT(	GGT	ra Tr	CTAA GATT	TGT ACA	AGC ICG	CTG(	TAT; 4TA:	CAA GTT
18	R V			T	С	K *	A *	<u>\$</u>		<u>N</u>	<u>V</u>	G *	<u>T</u>	N *	<b>V.</b>	A *	W	Y	Q
181	CAGAA	ACCA	G (	GGCA	ATC	TCC	TAA	AGC	ACTG	ATT	OAT?	CTCC	GT	CATC	CTA	CCG	GTAG	CAG1	rgga VCCT
38	GTCTT Q K		C ( G	CCGT' O		AGG P	ATT			I	Y	مهر ع	_ <u>S</u> _	SIAG	Y	R	Y	S	G
30	ν	-	•									A	*	* C	DR	# #2	☆	å	
241	GTCCC CAGGG	TGAT	C	GCTT	CAC	AGG	CAG'	TGG:	ATCT	GGG	SAC	AGA!	TT A A	TCAC	TCT	CAC	CAT(	CAGO	CCAT
58	V P		R	CGAA F	T T	G	S	G	S	G	T	D	F	т	L	T	I	S	Н
201	GTGCA	CULCU	YC.	A A C A	Մար	CCC	AGA	СТА'	TTTC	TG:	rca	GCA	ΑТ	ATAA	CAT	СТА	TCC'	rct	CACG
301	CACGT	CAGA	C	TTCT	GAA	.CCG	TCT	GAT.	AAAG	AC	AGT	CGT'	TA	TATI	GTA	GAT	AGG	AGA	GTGC
78	V Q	S	E	D	L	A	D	Y	F	С	Q *	Q *	<u>x</u>	N_	_ <u>I_</u>	* - <u>X</u>	<u>p</u>	<u>.l.</u>	<u>-</u> ↓
															CDR	#3			
361	TTCGG	יחירטיוו	c	ccac	CAA	.GCT	GGA	B GCT	stBI TCGA	AG	AGC	TGT	GG	CTGC	CACC	ATC	TGT	CTT	CATC
301	AAGCC	AGGA	C	CCTG	GTI	'CGA	CCT	CGA	AGCT	TC	TCG	ACA	CC	GACC	FTGG	TAG	ACA:	GAA	GIAG
,	F G	P	G	Т	K	L	E	L	R			V		A		_	•	_	
421	TTCCC	GCCA	T	CTGA	TGA	GCA	GTT	GAA	ATCT	GG.	AAC	TGC	TT	CTGT	rtgi	GTG	CCT	GCT(	GAAT CTTA
118	AAGGG	P P	S'A	GACT D	E E	Q	L	K	S	G	T	A	S	V	V	С	L	L	N
481	AACTI TTGAA	CTAT	rc C	CCAG	AGA	AGGC PCCG	CAA	AGT TCA	'ACAG TGTC	TG AC	GAA CTT	GGT CCA	SS.	ATAI	ACGC	CCT GGA	CCA GGT	ATC TAG	GGGT CCCA
138	N F	Y	P	R	E	A	K	V	Q	W	K	v	D	N	A	L	Q	S	G
541	AACTO	CCA(	3G	AGAG	TGT	CAC	AGA	GCA CGT	GGAC	AG	CAA	GGA	CA GT	GCA	CCTA GGAT	ACAG TGTC	CCT GGA	'CAG GTC	CAGC GTCG
158	N S	Q	E	S	v	T	E	Q	D	S	K	D	S	T	Y	S	L	S	S
601	ACCCT	rGAC	GC	TGAC	CA	AAGC	AGA	CTA	CGAG	AA .	ACA	CAA	LAG	TCT	ACG(	CTG	CGA	AGT	CACC
178	TGGG! T L	ACTG( T	CG L	ACTO	GT". K	TTCG A	TC1	Y Y	E E	K	H H	K	V	Y	A	C	E	V	T
661	CATC	AGGG(	CC	TGAC	SCT(	CGCC	CGI	CAC	CAAAC Crerence	AC	CTI	ኒር <u>አ</u> ያ	\CA Ст	GGG	GAGZ CTC'	AGTO	<b>:</b>		
198	H Q	G	L	S	S	P	V	T	K	S	F	N	R	G	E	C			
	L TT	TT										F	G	ì. 1	9				
21/		}																	

SUBSTITUTE SHEET (RULE 26)

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					ATAT			man	merii.	maa x	men	מו א מו	- ՄԱՄԻՆ	<b>-</b> C	արարարդու	יייייי	ייעמי	יויכריו	מחמי	ממ.
1	ATG	ለሊላ ጥጥባ	AAG TTTC	SA T	TATA	CGC/ SCG'	PAA TAA	AGA	AGA.	ACGT	AG	ATA	CAA	GC	AAAA	AAG	ATA	ACGA	TGI	TTG
-23				N	I	A	F		L		S	M	F		F	S	I	A	T	N
61	GCG	TAC	:GCI	G	AGGT	GCA	<b>SCT</b>	GGT	GGA	GTCT	GGC	GG.	AGG	CT	TAGT	GCC	GCC	TGG	\GGC	TCC
					TCCA						CCC	CCC,	rcc		ATCA V	CGG( P	CGG P	ACC'	rccc G	:AGG S
-3	A	Y	A	E	V	Q	L	V	E	S	G	G	G	L	V	P	r	G	G	3
121	CTG	AAA	CTC	T	CCTG	TGC	AGC	CTC	TGG	ATTC	ATA	ATT	CAG'	TA	GTTA'	TGG	CAT	GTC	TGC	GTT
	GAC	TTT	GAC	SA	GGAC.	ACG'	rcg	GAG	ACC	TAAG	TA	<b>CAA</b> 1	GTC	AΤ	CAAT.	ACC	GTA	CAG	ACC	CAA
18	L	K	L	S	С	A	A	S	G_	<u> F</u>	I_	F_	_S	_ <u>S_</u>	<del></del> X	G	M	S *	W	V
												<u></u>	DR :	#1	-	•	•	-		
												<u> </u>		₩ #						
181	CGC	CAC	ACI	C	CAGG	CAA	GAG	CCT	GGA	GTTG	GT	CGC.	AAC	CA	TTAA	TAA'	TAA	TGG	rgan	AGC
	GCG	GTC	TGA	λG	GTCC	GTT					CAG	3CG								
38	R	Q	$\mathbf{T}$	P	G	K	S	L	E	L	V	A	T	I	И	<u>N</u> _	_N	<u>G</u>	_D	S
													*	*	*	*	*	*	*	Ħ
2/1	ACC.	יייםיי	יתיםיו	יר	CAGA	CAG	TGT	GAA	GGG	CCGA	TT	CAC	CAT	CT	CCCG	AGA	CAA	TGC	CAAC	SAAC
741	TGG	AT	LTA	4G	GTCT	GTC	ACA	CTT	CCC	GGCT	AA	GTG	GTA	GA	GGGC	TCT	GTT	ACG	GTTC	TTG
58	T	Y	Y	P	D	S	V	K	G	R	F	T	I	S	R	D	И	A	K	N
	*	*	* CDR	# <b>-</b>	*	*	*	*					-							
•																			•	
301	ACC	CTC	3TAC	CC	TGCA	TAA	GAG	CAG	TCT	GAAG	TC	TGA	GGA	CA	CAGC	CAT	GTT	TTA	CTG:	rgca
					ACGT						AG. S		CCT D	GT T	GTCG A	GTA M	F	AAT Y	C	A
78	T	L	Y	L	. <b>Q</b>	M	S	S	L	V	3	E.	D	1	A	**	•	•,	•	••
361	AGA	\GC(	CTC	CA	TTAG	TTC	GGC	TAC	TTG	GTTT	GG	тта	CTG	GG	GCCA	AGG	GAC	TCT	GGT	CACT
	TCT	CGC	GGA(	ЗT	AATC	AAG	CCG	ATC	AAC	CAAA	CC	AAT	GAC	CC	CGGT	TCC	CTG	AGA	CCA	FIGA
98	R	Α	<u>L</u>	I	S	_S.	<u>A</u>	T		F_	G	Y	W	G	Q	G	T	Ь	V	T
		*	*	*	*	*		*	*	*	*	*								
						CD	R #	3 Apa	. т											
A 2 1	CTC	ישרי	rcc	ΔC	CCTC	יראר	מבחי	GGC	CCC	ATCG	GT	CTI	ccc	:cc	TGGC	ACC	CTC	CTC	CAA	GAGC
	CNC	200	אכפי	TC.	CCAC	CTC	ርጥጥ	CCC	GGG	TAGC	CA	GAA	GGG	GG	ACCG	TGG	GAG	GAG	GTT	CTCG
118	v	S	Α	A	S	T	K	G	P	S	V	F	P	L	A	P	S	S	K	S
481	ACC	CTC'	TGG	GG	GCAC	CAGC	GGC	CC	rGGC	CTGC	CI	CC2	CAA	CC	TGAT	rgaa	GGG	GCT	TGG	CCAC
138	TGG	JAG. S	ACC G	CC G	T	A	A	L	G	C	L	v	K	D	Y	F	P	E	P	V
541	ACC	GGT	GTC	GT	GGAZ	ACTO	AGG	CGG		rgacc	AG	CGC	CC	rgc	ACAC	CTI	יכככ אכפפ		ACA	GGAT
150	TG(	CCA	CAG	CA ผ	CCT	rGA(	TCC G	. GC(	افاقاد ۲۰	なこまらし グ	, IC	.GCC	V	ico H	T	F	.000 P	A	v	L
601	CA	GTC	CTC	AG	GAC	CT	ACTO	CC'	rca	GCAGC	: G1	GG?	rgac	CCG	TGC	CTC	CAG	CAC	CTI	GGGC
<b>.</b>	GT	CAG	GAG	TC	CTG	AGA'	r <b>GA</b> G	GG	AGT	CGTCC	CA	YCC!	ACTO	GC u	ACG(	)Aن ح	SGTC C	GTC C	.ска Гі	G
178	Q	S	S	G	L	Y	S	L						V	P	3	3	J	•	_
									F	IG	9	n	Δ		•					

## **FIG. 20A**

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- 661 ACCCAGACCT ACATCTGCAA CGTGAATCAC AAGCCCAGCA ACACCAAGGT GGACAAGAAA TGGGTCTGGA TGTAGACGTT GCACTTAGTG TTCGGGTCGT TGTGGTTCCA CCTGTTCTTT 198 T Q T Y I C N V N H K P S N T K V D K K
- 721 GTTGAGCCCA AATCTTGTGA CAAAACTCAC ACATGA CAACTCGGGT TTAGAACACT GTTTTGAGTG TGTACT

K T H218 V E P K S C D

FIG. 20B

Light Cl	nain Primers:	
MKLC-1,	22mer	
-5 '	CAGTCCAACTGTTCAGGACGCC 3'	
MKLC-2,	22mer	
5'	GTGCTGCTCATGCTGTAGGTGC 3'	
MKLC-3,	23mer	
5 '	GAAGTTGATGTCTTGTGAGTGGC	3 '
	hain Primers: 1, 24mer	
5'	GCATCCTAGAGTCACCGAGGAGCC	3 '
IGG2AC-	2, 22mer	
5 '	CACTGGCTCAGGGAAATAACCC 3'	
IGG2AC-	3, 22mer	
5'	GGAGAGCTGGGAAGGTGTGCAC 3'	
	FIG. 21	

6G4.light.Nsi 36-MER

5' CCAATGCATACGCT GAC ATC GTG ATG ACC CAG ACC CC 3'

T T T T T

A A

Light chain reverse primer

6G4.light.Mun 35-MER

5' AGA TGT CAA TTG CTC ACT GGA TGG TGG GAA GAT GG 3'

Light chain forward primer

FIG. 22

FIG. 23

1.338.299

70 G ATATCGTGAT GACACAGACA CCACTCTCCC TGCCTGTCAG TCTTGGAGAT C TATAGCACTA CTGTGTCTGT GGTGAGAGGG ACGGACAGTC AGAACCTCTA 1 D I V M T Q T P L S L P V S L G D 121 CAGGCCTCCA TCTCTTGCAG ATCTAGTCAG AGCCTTGTAC ACGGTATTGG AAACACCTAT GTCCGGAGGT AGAGAACGTC TAGATCAGTC TCGGAACATG TGCCATAACC TTTGTGGATA 18 Q A S I S C R S <u>S O S L V H G I G N T Y</u> \* \* \* \* \* \* \* CDR #1 181 TTACATTGGT ACCTGCAGAA GCCAGGCCAG TCTCCAAAGC TCCTGATCTA CAAAGTTTCC AATGTAACCA TGGACGTCTT CGGTCCGGTC AGAGGTTTCG AGGACTAGAT GTTTCAAAGG 38 L H W Y L Q K P G Q S P K L L I Y CDR #2 241 AACCGATTTT CTGGGGTCCC AGACAGGTTC AGTGGCAGTG GATCAGGGAC AGATTTCACA TTGGCTAAAA GACCCCAGGG TCTGTCCAAG TCACCGTCAC CTAGTCCCTG TCTAAAGTGT 58 N R F S G V P D R F S G S G S G T D F T 301 CTCAGGATCA GCAGAGTGGA GGCTGAGGAT CTGGGACTTT ATTTCTGCTC TCAAAGTACA GAGTCCTAGT CGTCTCACCT CCGACTCCTA GACCCTGAAA TAAAGACGAG AGTTTCATGT 78 L R I S R V E A E D L G L Y F C S CDR #3 361 CATGTTCCGC TCACGTTCGG TGCTGGGACC AAGCTGGAGC TGAAACGGGC TGATGCTGCA GTACAAGGCG AGTGCAAGCC ACGACCCTGG TTCGACCTCG ACTTTGCCCG ACTACGACGT 98 H V P L T F G A G T K L E L K R A D A A MunI 421 CCAACTGTAT CCATCTTCCC ACCATCCAGT GAGCAATTGA GGTTGACATA GGTAGAAGGG TGGTAGGTCA CTCGTTAACT 118 P T V S I F P P S S E Q L K FIG. 24

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70 G AGATTCAGCT GCAGCAGTCT GGACCTGAGC TGATGAAGCC TGGGGCTTCA C TCTAAGTCGA CGTCGTCAGA CCTGGACTCG ACTACTTCGG ACCCCGAAGT G P E L M K P Q Q S I O L 121 GTGAAGATAT CCTGCAAGGC TTCTGGTTAT TCATTCAGTA GCCACTACAT GCACTGGGTG CACTTCTATA GGACGTTCCG AAGACCAATA AGTAAGTCAT CGGTGATGTA CGTGACCCAC 18 V K I S C K A S G Y S F S S H W V \_H Y M CDR #1 181 AAGCAGAGCC ATGGAAAGAG CCTTGAGTGG ATTGGCTACA TTGATCCTTC CAATGGTGAA TTCGTCTCGG TACCTTTCTC GGAACTCACC TAACCGATGT AACTAGGAAG GTTACCACTT I G Y I 38 K Q S H G K S L E W **CDR #2** 241 ACTACTTACA ACCAGAAATT CAAGGGCAAG GCCACATTGA CTGTAGACAC ATCTTCCAGC TGATGAATGT TGGTCTTTAA GTTCCCGTTC CGGTGTAACT GACATCTGTG TAGAAGGTCG Q K F K G K A T L T V D T 301 ACAGCCAACG TGCATCTCAG CAGCCTGACA TCTGATGACT CTGCAGTCTA TTTCTGTGCA TGTCGGTTGC ACGTAGAGTC GTCGGACTGT AGACTACTGA GACGTCAGAT AAAGACACGT H L S S L T S D D S A V Y F C A 78 T A N V 361 AGAGGGGACT ATAGATACAA CGGCGACTGG TTTTTCGATG TCTGGGGCGC AGGGACCACG TCTCCCCTGA TATCTATGTT GCCGCTGACC AAAAAGCTAC AGACCCCGCG TCCCTGGTGC 98 R G D Y R Y N G D W F F D V W G A G T T \* \* \* **CDR #3** BstEII 421 GTCACCGTCT CCTCCGCCAA AACCGACAGC CCCATCGGTC TATCCGGGCC CAGTGOCAGA GGAGGCGGAT TTGGCTGTCG GGGTAGCCAG ATAGGCCCGG PIGL 118 V T V S S A K T D S 471 CATC GTAG 135 I

FIG. 25

5' CTTGGTGGAGGCGGAGACG 3'

Mutagenesis Primer for 6G425VL

DS/VF 38MER

5' GAAACGGGCTGTTGCTGCACCAACTGTATTCATCTTCC 3'

SYN.BstEII 31 MER

5' GTCACCGTCT CCTCCGCCTC CACCAAGGGC C 3'

SYN.Apa 22 MER

5' CTTGGTGGAGGCGGAGGACG 3'

FIG. 26

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1	ATGA TACT	AGA. TCT	AGA TCT	TATA	GCG	TAA	AGA	AGA	ACGT	AG!	)ATA	CAAC	3C	TTTTI	AAG	ATA	ACGA	TGT	TTA
-23	M K	K	N	I	A	F	L	L	A	S	M	F	V	F	S	I	. <b>A</b>	T	N
61	GCAT CGTA	ACG TGC	CTG GAC	TATA	GCA	CTA	CTG'	rgt	CTGT	GG'	rgac	GAGC	3G	TGCC'	ACAC	STC	AGAA	CCT	CTA
	A Y			_	V			Q						P			L		_
	GTCC	GGA	GGT	AGAG	AAC	GTC	TAG	ATC.	AGTC	TCC	3GA/	CAT	ľG	ACGG'	<b>LATA</b>	ACC	TTTC	CACC STGG T	TAT ATA
18	Q A	S	I	S	С	R ⋆	S ★	*		*	* CDR	*	*	<u> </u>	*	*	*	*	*
	AATG	TAA	CCA	TGGA	CGT	CTT	CGG	TCC	GGTC	AG	AGG'	rtt(	CG	TCCT AGGA L	CTA	<b>GAT</b>	GTT	AGTT CAA V	AGG
	* *																* CDR	#2	*
241	AACC	GAT CTA	TTT AAA	GACC	CCA	GGG	TCT	GTC	CAAG	TC.	ACC	GTC	AC	GATC CTAG	TCC	CTG	TCT	AAAC	ACA TGT
58	N F	-	* S	G	V	P	D	R	F	S	G	S	G	S	G	T	D	F	T
301	CTCA	GGA	TCA	GCAC	AGT	GGA	GGC	TGA	GGAT	CT	GGG.	ACT'	TT	ATTT TAAA	CTG GAC	CTC	TCA.	AAGT TTC	ACA ATGT
78				R			A	E	D	L	G	L	Y	F	С	S	Q	<u>s</u>	$\underline{\mathbf{T}}$
							÷					-					OR#	3	
361	CATO	TTC AAC	CGC GCG	TCAC AGTC	GTI SCA	4GCC	ACG	ACC	CTGG	$\mathbf{T}\mathbf{T}$	CGA	CCT	CG	TGAA	'TGC	CCG	ACA	ACG	ACGT.
98	<u>H</u>	7 -	<u> </u>	T *	F	G	A	G	Т	K	L	E	L	Κ.	R	Α	V	A	A
	CCT	יכאכ	מדמי	AGT	AGA	AGGG	TGC	STAC	GTCA	CT	CGT	TAA	CT	AATC	BACC	TTG	ACG	GAG	ACAA
118	P !	r 1	V F	· I	F	P	P	S	S	E	Q	L	K	<i>S</i>	G	T	A	S	V
	CAC	ACGO	BACG	ACT'	TAT	<b>rgaa</b>	GA'	CAGC	GTCT	CI	CCG	GTT	TC	TACA	CAC	CTT	CCA	CCT	ATTG
														Q					
	CCC	SAGO	ፈጥጥል	GCC	CAT	TGAG	GG?	rcc:	<b>CTCA</b>	CA	GTC	TCT	rcg	AGG	rgro	GTT	CCI	GTC	GIGG
														D					
	አጥር	TCC	മരന	י ככת	CGT	GGGA	CTO	GCG	ACTCC	3 T7	rtcc	TCI	rga	ACG	rcr:	rtgt	' GT"	"I'CA	GATG
178	3 Y	S.	L S	s s	T	L	T	L	s	K	A	D	Y	T E	K	H	K	V	Y

# FIG. 27A

661 GCCTGCGAAG TCACCCATCA GGGCCTGAGC TCGCCCGTCA CAAAGAGCTT CAACAGGGAA CGGACGCTTC AGTGGGTAGT CCCGGACTCG AGCGGGCAGT GTTTCTCGAA GTTGTCCCCT 198 A C E V T H Q G L S S P V T K S F N R G

-1G. 27B

CTCACAATT E C O

218

721 GAGTGTTAA

1	ATGA	AAZ	\AG/	<b>.</b> .	ATATO	:GC	\TT	TCT	TCT	rgca	TCI	YTA'	TTC	:G	TTTT	rrci	TAT	TGC:	rac:	AAAC	!
_	TACT	TT.	rrc	ר י	TATAC	GCG	AAT	AGA	AGA	ACGT	AG/	YAC	CAAC	C:	AAAA	AAGA	ATA	ACG	ATG'	LLIG	;
	M K	: 1	K 1	1	I	A	F	L	L	A	S	M	F	V	F	S	I	A	T	'N	
61	GCGT	'AC	CTC	3 7	AGAT?	rca(	GCT CGA	GCA	GCA(	GTCT CAGA	GGZ CCT	CCI CGG	GAC ACTO	SC CG	TGAT(	GAA( CTT(	GGG	TGG(	GGC'	ITCA AAGI	•
	A Y	1	A I	Ε	I	Q	L	Q	Q	S	G	P	E	L	M	K	P	G	Α.	S	
121	GTGA	AG	ATA!	r (	CCTG(	CAAC STTC	GCC CCG	TTC AAG	TGG ACC	TATT AATA	TCA AG	TTE AA1	CAGI	A? T/	GCCA(	CTA( GAT(	CAT GTA	GCA(	CTG GAC	GGTG CCAC	}
18	V K						A	S	G_	_Y	S	F	S	<u>s</u>	<u>H</u>	Y *	M *	H *	W	V	
													CDI								
181	AAGC	CAG	AGC	C .	ATGG	AAA	GAG	CCT	TGA	GTGG	AT'	rgg	CTAC	CA	TTGA	TCC'	TTC	CAA	TGG	TGAZ	<b>\</b>
							CTC	GGA	ACT	W	TA	G		I	AACT.	P_	S	N	G	E.	•
38	K Ç	)	S	Н	G	K	S	Ъ	E	w	1	G	*	*	*	. <del>*</del>	*	*	*	*	
															C	DR	#2				
241	ACT/	ACT IGA	TAC.	A T	ACCA TGGT	GAA CTT	ATT TAA	CAA	.GGG	CAAG	GC	CAC.	YTTA AAT	GA CT	CTGT GACA	AGA TCT	CAC GTG	ATC TAG	TTC AAG	CAG(	3
58		י ני		N	Q		F	K	G	K	A	T	L	$\mathbf{T}$	v	D	T	S	S	S	
	* 1	k	*	*	*	*	*	*	*												
301	ACAC	GCC	AAC	G	TGCA	TCT	CAG	CAC	CCI	GACA	TC	TGA	TGA	CT	CTGC GACG	AGT	CTA	TTT	CTC	TGC.	A T
										T		MC I	D	2	A	v	Y	F	C	Α	_
					Н													_			_
361	AGA	GGG	GAC	$\mathbf{T}$	ATAG	ATA	CAA	CGG	CGA	CTGG	TT	TTT	CGA	TG	TCTG	GGG	CGC	AGG	GAC	CAC	G
															AGAC	:CCC	GCG	TCC	T.	T'	U
98	R (	3	D	Y_	R	_Y_	_N_				<u> </u>		_D	V	W	G	A	G	T	1	
	•	*	*	*	*	¢ CE	* R #	3	*	*	*	*	*	*							
421	GTC	ACC	GTC	T	CCTC	CGC	CTC	CAC	CAI	AGGGC	CC	ATC	GGT	CT GA	TCCC AGGC	CCT	GGC CCG	ACC	CTC	CCTC SGAG	C G
118						A		T	K	G	P	S	V	F	P	L	A	P	S	S	
481	AAG.	AGC	ACC	T:	CTGG	GGG	CAC	AG(	CGG(	CCTG	GG	CTC	CCT GGA	SS'	TCA	AGGA rcci	CTA GAT	CTT GA	rcc( Agg(	CCGA GCT	A T
138	K	s	T	S	G	G	T	A	A	L	G	C	L	ν	K	D	Y	F	P	E	
	GGC	CAC	TGC	C	ACAG	CAC	CTI	GA	GTC	CGCGG	GA	CTC	GTC	:GC	GCG7	ACG7	rgtg	GA	AGG	GCCG	T A
	P	V	T	V	. <i>S</i>	W	N	S	G	A	L	T	S	G	·	H	T	F	P	A	
	CAG	GAT	rgre	'A	GGAC	TCC	CTGA	GA'	TGA	GGGAC	TC	GTC	CGCA	CC	TGAC	GGC)	<b>ACG</b> G	GA	GGT	CGTC	SC CG
178	V	L	Q	S	S	G	L	Y	S	L	S	S	v	V	T	V	P	S	S	S	

# FIG. 28A

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>

GGTCGTTGTG z

TTAGTGTTCG

GACGTTGCAC

Y T

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198

Ŋ 721 AAGAAAGTTG AGCCCAAATC TTGTGACAAA ACTCACACAT GA TTCTTTCAAC TCGGGTTTAG AACACTGTTT TGAGTGTGTA CT H Ħ Ħ TICTITICAAC TCGGGTTTAG AACACTGTTT D ن ا AACCCGTGGG TCTGGATGTA

×

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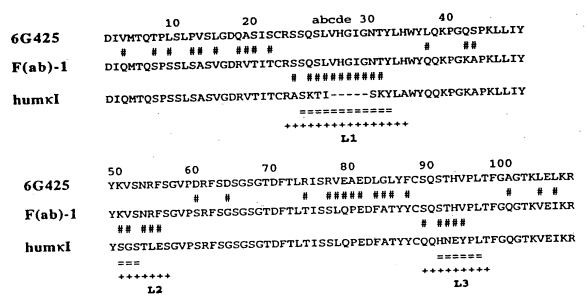
×

218

661 TTGGGCACCC AGACCTACAT CTGCAACGTG AATCACAAGC CCAGCAACAC CAAGGTGGAC

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#### Variable Light Chain Domain



## Variable Heavy Chain Domain

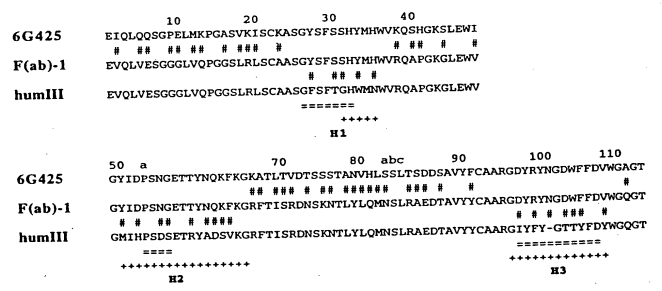
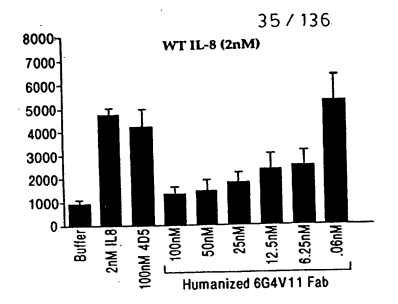


FIG. 29



**FIG. 30A** 

IC50~12nM

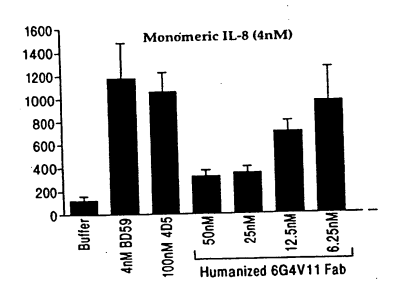


FIG. 30B

IC50~15nM

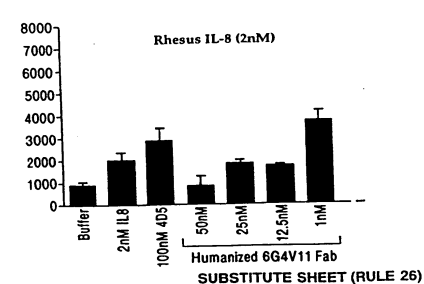


FIG. 30C

IC50~22nM

anti-IL-8 6G4.2.5V11 Light Chain Amino Acid Sequence of the humanized

ALQSGNSQESVTEQDSKDSTYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRG LHWYQQKPGKAPKLLIYKVSNRFSGVPSRFSGSGSGTDFTLTISSLQPEDFATYYCSQST HVPLTFGQGTKVEIKRTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDN MKKNIAFLLASMFVFSIATNAYADIQMTQSPSSLSASVGDRVTITCRSSQSLVHGIGNTY

anti-IL-8 6G4.2.5V11 Heavy Chain Amino Acid Sequence of the humanized

WVRQAPGKGLEWVGYIDPSNGETTYNQKFKGRFT**L**SRDNSKNT**A**YLQMNSLRAEDTAVYY CARGDYRYNGDWFFDVWGQGTLVTVSSASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTK MKKNIAFLLASMFVFSIATNAYAEVQLV**Q**SGGGLVQPGGSLRLSCAASGYSFSSHYMH VDKKVEPKSCDKTHT Amino Acid Sequence of the peptide linker and M13 Phage Coat (gene-III)

GLANGNGATGDFAGSSNSQMAQVGDGDNSPLMNNFRQYLPSLPQSVECRPFVFSAGKPY SGGGSGSGDFDYEKMANANKGAMTENADENALQSDAKGKLDSVATDYGAAIDGFIGDVS EFSIDCDKINLFRGVFAFLLYVATFMYVFSTFANILRNKES

# FIG. 31A

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									•							~~~		maa.	n» a :	
1	ATG	AAA	AAC	SA.	ATAT	CGC.	ATT	TCT'	rct'	rgca	TCI	'ATG	TTC	G'	TTTTT	PTCT	MT.	TGC:	IAC	MAAC
	TAC'	TTT	TTC												AAAA					
-23	M :	K	K	N	I	Α	F	L	L	A	S	M	F	V	F	S	I	Α	Т	N
61	GCA'	TAC	GCI	rg	TATA	CCA	GAT	GAC	CCA	GTCC	CCC	SAGC	TCC	C '	TGTC	CGCC	rrc	TGT	GGG(	CGA'I'
	CGT	ATG	CGA	AC	TATA	GGT	CTA	CTG	GGT	CAGG	GGC	CTCC	SAGG	G.	ACAG	GCGG	SAG	ACA	CCC	GCTA
-3	Α	Y	Α	D	I	Q	M	$\mathbf{T}$	Q	S	P	S	S	L	S	A	S	V	G	D
													•							
121	AGG	GTC	ACC	CA	TCAC	CTG	CAG	GTC.	AAG'	TCAA	AGO	CTT	\GTA	/C	ATGG'	TAT	\GG	TAA	CAC	GTAT
	TCC	CAG	TG	ЗT	AGTG	GAC	GTC	CAG	TTC.	AGTT	TCC	GAAT	CAT	rG	TACC.	ATA	rcc	ACG.	ATG	CATA
18	R	v	${f T}$	I	T	С	R	S	S	Q	S	L	V	H	G	I	G	N	T	Y
181	TTA	CAC	TG	GT	ATCA	ACA	GAA	ACC	AGG	AAAA	GC'	rcco	SAAZ	AC.	TACT	GAT'	ГТА	CAA	AGT.	ATCC
	AAT	GTO	SAC	CA	TAGT	TGI	CTT	TGG	TCC	TTTT	CG	AGG	CTT	rg	ATGA	CTA	TAA	GTT	TCA	TAGG
38	L	Н	W	Y	0	Q	K	P	G	K	A	P	K	L	L	I	Y	K	V	S
241	TAA	rcg/	TT	СТ	CTGG	AGT	rccc	TTC	TCG	CTTC	TC	TGG	ATC	CG	GTTC	TGG	GAC	GGA	TTT	CACT
211	TTA	GC	יאמי	GA	GACC	TCA	AGGG	AAG	AGC	GAAG	AG.	ACC'	TAG	GC	CAAG	ACC	CTG	CCT	'AAA	GTGA
5.8	N	R	F	S	G	v	P	S	R	F	s	G	S	G	S	G	${f T}$	D	F	T
50	••	•	•		Ū	•	-													
301	CTC	200	יים י	CA	GCAC	ישרי	rgca	GCC	:AGA	AGAC	TT	CGC.	AAC'	$\mathbf{T}\mathbf{T}$	ATTA	CTG	TTC	ACA	GAG	TACT
301	CIC	ייייני דייייי	מחב	CT.	CGTC	TAG	ACGT	CGG	TCT	TCTG	AA	GCG	TTG.	AA	TAAT	GAC	AAG	TGI	CTC	ATGA
70	L					T.	0	P	E	D	F	Α	Т	Y	Y	С	S	Q	S	T
70	ь		_	5		~	×	•	_											
261	CAT	rcm			TCAC	برات	ኮጥርር	ACZ	AGGG	TACC	. AA	GGT	GGA	GA	TCAA	ACG	AAC	TGT	rGGC	TGCA
201	CM	IGI		CC.	ACT(	COL	2200	TG	יייייי	ATGG	тт	CCA	CCT	CT	AGTI	TGC	TTG	ACA	ACCG	ACGT
0.0					AGI	JCA.	nncc C	10.		т	ĸ	v	E	I	K	R	${f T}$	v	Α	A
98	Н	V	P	1	1	г	G	Q	G	•	••	·	_							
401	~~	3 mc	men		mc a t	ጥርጥ	ጥሮርር	CCC	ጉ ል ጥር	רתכאח	r GA	GCA	GTT	ĠΑ	AATO	CTGG	AAC	TG	CTTC	TGTT
421	CC	MAC	IGI	(C)	1CA	1C1	ACCC	. GC(	ጋጥ አ <i>(</i>	77.01.1 77.04.5	ייס א	CGI	CAA	CT	TTAC	SACC	TTG	AC	GAAC	BACAA
110						AUA T	AGGC	יטטי ס	SIA	D	. C.	0	L	ĸ	s	G	Т	Α	S	٧.
118	P	5	V	r	1	F			J		_	×	_							
401		~m~		ncc	mc x	ת יווו ת	አርሞባ	י כייי	מיים מ	CCAG	A GE	AGGC	CAA	AG	TAC	AGTO	GAA	. GG	rggi	ATAAC
481	GT	GIG	CCI	rGC	TGA	WIW		CA	MAC.	CCMC	ייטי ייטיי	ייייייי	ייייטי	ייויר	ATG	TCAC	CTT	· CC	ACC'	PATTG
						TAT	TGAF	. GA	TAG	9 TC.	r C.	אכטנ	W.	v	Q	w	ĸ	v	D	N
138	V	C	ь	L	, N	N	ır	1	P	K	E	А	10	٠	×	•		-		
											n C	ריי א ני	~~~	\CC	) AGG	מרמנ	CAR	. GG	ACA	GCACC
541	. GC	CCI	CC	LAA	' CGG	GTA	ACT	2 00	AGG.	MOMO	i G		าเมตะ วาเมตะ	rcc	ייייייייייייייייייייייייייייייייייייייי	TCT(	יכתים יכתים		TGT	CGTGG
	CG	GG?	\GG'	TTA	GCC	CAT	TGA	3 GG	TCC	TCTC	A C/	AGT(	21C1		100	1010	.GI.	. כס	202	CGTGG T
158	3 A	L	Q	S	; G	; N	1 S	Q	E	. S	V	Т	£	Ų	D	3	K	ט	٥	-
																			አአር	ጥርጥልር
601	L TA	CAC	GCC'	TC	A GCA	\GC#	ACCC.	r ga	CGC	TGAG	C A	AAG	CAGA	ACI	ACG	AGA		CO	MMC	TCTAC
	ra	CTO	CGG.	AGI	r CG1	rcgi	rggg	A CI	'GCG	ACTC	G T	TTC	GTC'	I'GA	1 TGC	TCT	1"I'G'.	r Gi	110	AGATG
178	B Y	s	L	5	5 5	3	r L	T	L	, S	K	Α	D	Y	E	K	н	, r	. •	1
																				CCCCA
663	1 GC	CTC	GCG	AAC	TCA	ACC	CATC	A GC	GCC	TGAG	СТ	CGC	CCG'	TC	A CAA	AGA	GCT".	r CA	ACA	GGGGA
	CC	267	$\neg cc$	ጥጥር	AG	rcc	TAG	T CC	CGG	ACTC	GA	GCG	GGC	AGT	r GTT	TCT	CGA	A G1	1.61	CCCCI
19	8 A	C	E	, ,	J :	r 1	H Q	C	3 I	S	S	P	V	7	r K	S	F	V	ı R	G
72	1 G	AGT	GTT	'AA	G CT	GAT	CCTC	T AC	CGCC	CGGAC	G C	ATC	GTG	GC	CTA	GTA	CGC	A AC	TAC	TCGTA
	C	rca	CAA	TT	C GA	CTA	GGAG	A TO	GCGC	CCTC	CG	TAG	CAC	CG	G GAI	CAT	GCG	T TC	SATC	AGCAT
21	8 E																	•		
									F	— n ∕	_		</td <td>(( )</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>	(( )						

FIG. 31B

anti-IL-8 6G4.2.5V19 Light Chain Amino Acid Sequence of the humanized

ALQSGNSQESVTEQDSKDSTYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRG LHWYQQKPGKAPKLLIYKVSNRFSGVPSRFSGSGSGTDFTLTISSLQPEDFATYYCSQST HVPLTFGQGTKVEIKRTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDN MKKNIAFLLASMFVFSIATNAYADIQMTQSPSSLSASVGDRVTITCRSSQSLVHGIGNTY

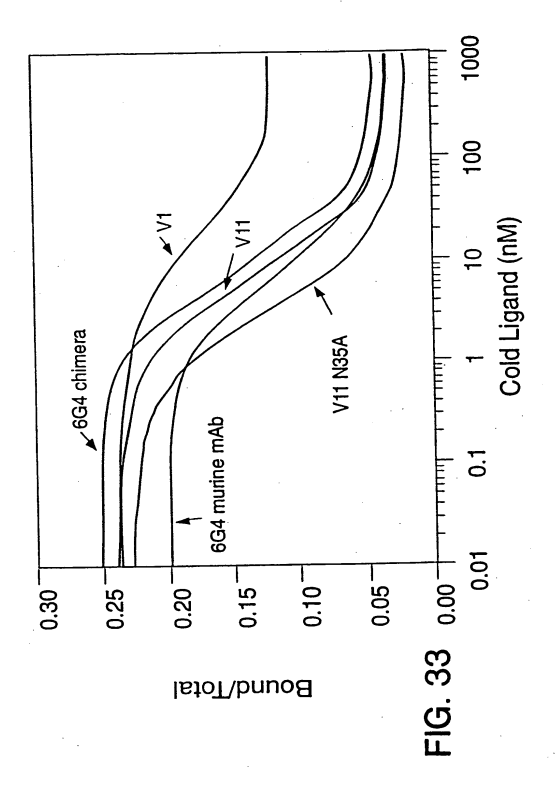
anti-IL-8 6G4.2.5V19 Heavy Chain Amino Acid Sequence of the humanized

WVKQAPGKGLEWVGYIDPSNGETTYNQKFKGRFTLSRDNSKNTAYLQMNSLRAEDTAVYY CARGDYRYNGDWFFDVWGQGTLVTVSSASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTK MKKNIAFLLASMFVFSIATNAYAEVQLVESGGGLVQPGGSLRLSCAASGYSFSSHYMH VDKKVEPKSCDKTHT

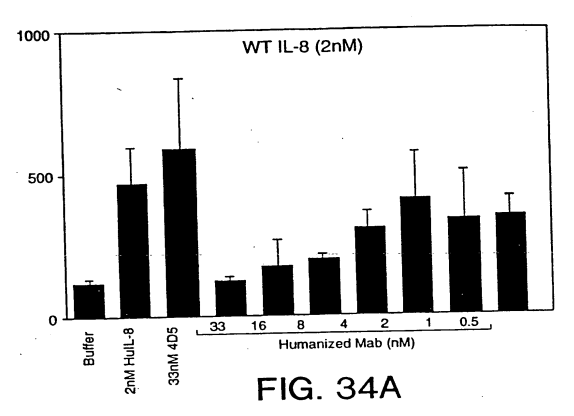
FIG. 31C



F16.32



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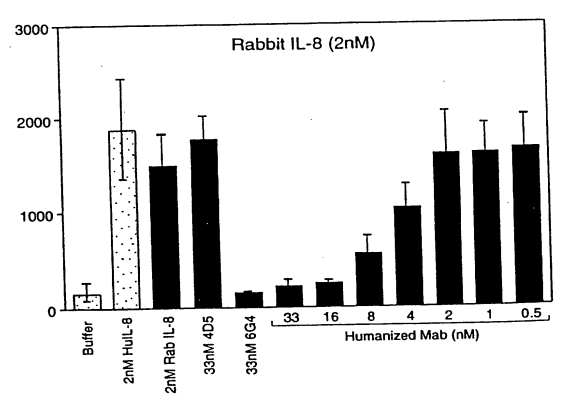
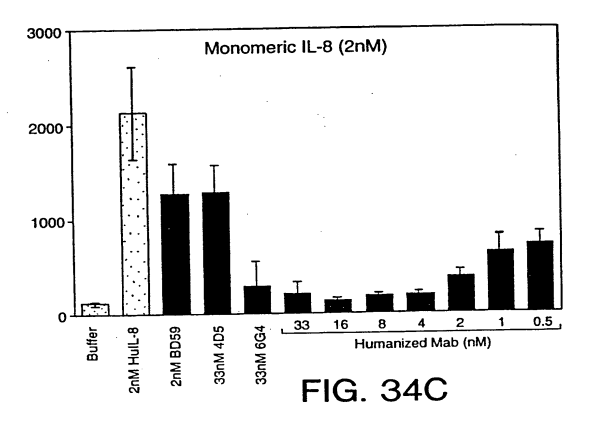
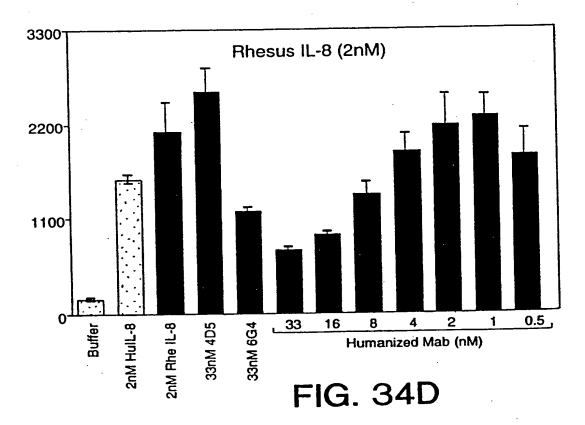


FIG. 34B

SUBSTITUTE SHEET (RULE 26)





SUBSTITUTE SHEET (RULE 26)

anti-IL-8 6G4.2.5V11N35A Light Chain Amino Acid Sequence of the humanized

ALQSGNSQESVTEQDSKDSTYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRG MKKNIAFLLASMFVFSIATNAYADIQMTQSPSSLSASVGDRVTITCRSSQSLVHGIG**A**TY LHWYQQKPGKAPKLLIYKVSNRFSGVPSRFSGSGSGTDFTLTISSLQPEDFATYYCSQST HVPLTFGQGTKVEIKRTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYPREAKVQWKVDN

Amino Acid Sequence of the humanized anti-IL-8 6G4.2.5V11N35A Heavy Chain

CARGDYRYNGDWFFDVWGQGTLVTVSSASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF WVRQAPGKGLEWVGYIDPSNGETTYNQKFKGRFTLSRDNSKNTAYLQMNSLRAEDTAVYY PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLSSVVTVPSSSSLGTQTYICNVNHKPSNTK MKKNIAFLLASMFVFSIATNAYAEVQLVQSGGGLVQPGGSLRLSCAASGYSFSSHYMH VDKKVEPKSCDKTHT Amino Acid Sequence of the putative Pepsin Cleavage Site and GCN4 Leucine Zipper

CPPCPAPE<u>LL</u>GGRMKQLEDKVEELLSKNYHLENEVARLKKLVGER

FIG. 35

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1	ATG	AAZ	AAG	ЗA	ATAT	CGC	ATT	TCT:	rcti	rgca	TC	TAT	TTC	CG	TTTT	rtci	TAT	TGCI	ACA	AAC
	TO A C	· TriTriTri	ישיי	THI.	ጥልጥልር	יברכי	ተል አ	AGA	<b>AGA</b>	ACGT	AG	\TAC	)AA:	GC	AAAA	AAGA	YTA .	ACGA	71G1	TIG
-23	M	K	K	N	I	A	F	L	L	A	S	M	F	V	F.	5	1	A	1	14
	000	1 m	700		ma ma		സമ	CTC	രസ	ገልርር	GGG	CTC	SAG	GG	TGTC	كالتالحا	JAG	MCM	-cc	2C IW
-3	A	Y	A	D	I	Q	M	T	Q	S	P	S	S	L	S	A	S	V	G	D
121	AGC	GT	CAC	CA	TCAC	CTG	CAG	GTC.	AAG'	rcaa	AG	CTT	AGT.	AC TG	ATGG TACC	TATA	AGG TCC	TGC:	rac(	GTAT CATA
18	TCC R	V	T'	GT	AGTG T	C	R_	S	S	0	s	L_	v_	H	G	I	G	_A_	T	<u>_</u> Y
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181	TT	ACA	CTG	GT	ATCA	ACA	GAA	ACC	AGG.	AAAA	GC	TCC	GAA	AC TC	TACT	CTA	חממ תממ	CAA	TCA'	TAGG
	AA'	rgt	GAC	CA	TAGT	TGT	'CTT	TGG	TCC	JIII.	CG.	AGG'	CTT	Į G	ATGA	T	v	K	v	S
															L					
241	AA'	TCG	ATT	СТ	CTGG	AGT	CCC	TTC	TCG	CTTC	TC	TGG	ATC	CG	GTTC	TGG	GAC	GGA'	TTT	CACT
	mm.	800	מ מיחי	CA	CACC	יתיים	CCC	AAG	:AGC	GAAG	AG	ACC	TAG	iGC.	CAAG	ACC	CIG	CCI.	WWW	GIGN
	<u>N</u>	R_	F	S	G	V	P	S	R	F	S	G	S	G	S	G	T	D	F	1
301	СТ	GAC	CAT	CA	GCAG	TCI	rgCA	GCC	AGA	AGAC	тт	CGC	AAC	TT	ATTA	CTG	TTC	ACA	GAG	TACT
	$C\lambda$	CTC	מיתים	CT	CCTC	יאכו	TOO	CGC	TCT	'TCTG	A.A	.GCG	TTG	jΑΑ	TAAT	JAJ.	AAG	161	CIC	AIGA
	L	T	I	S	S	L	Q	P	E	D	F	A	Т	Y	Y	C	5			<del></del>
361	CA	TGT	CCC	CGC	TCAC	GT?	rtgg	ACA	\GG@	TACC	AA	GGI	'GGA	AGA	TCA	AACG	AAC	TGT	GGC	TGCA
	~~		000	200	N COTO	7 C R 1	1 1 CC	സവ	רכככ	<b>ነ</b> ልጥርር	ויידי :	CCA	cc.	ľ	AGT		. 1 1 6	WC'U		ACGI
															K					
421	CC	'ATC	TG:	rct	TCA!	rct'	rccc	GC	CATO	TGAT	' G	AGC	GT	rga	AATO	CTGG	AAC	TGC	TTC	TGTT
		mac	1801	202	አርጥ	ACD!	ACCC	്രദ	TAC	ACTA	$\mathbf{C}$	regi	CA	ACT	TTA	JACL	.116	ACC		MCM.
	P	S	V	F	· I	F	P	P	S	D	E	Q	Ь	K	. 5	G	[1	A	3	•
481	GT	ነርጥ	GC'	TGC	TGA	ATA	ACTI	CT	ATC	CAGA	G	AGG	CCA	AAG	TAC	AGTO	<b>GAA</b>	GGT	rggz	ATAAC
	~ 1	03/	700	200	• እርጥ	ידי תידי	ጥርያል	CA	ፐልርር	GCTCI	r C'	$\mathbf{r}\mathbf{c}\mathbf{c}$	3GT"	T" $T$ $C$	AIG	LCW		CCA	100	171110
138	v	С	L	I	, N	N	F	Y	P	R	E	A	K	V	7 Q	W	K	V	Ð	N
F 4 1			ncc	2 2 1		<b>ር</b> ሞአ	አ ርጥር	٠	ACC:	AGAGI	r G'	TCA	CAG	AGC	AGG	ACA	GCAA	GGZ	ACAC	CACC
541			ICC.	mm i	CGG	CYT	TC IC	י ככ	TCC'	тстси	A C.	AGT	GTC'	TCC	TCC	TGT	CGTT	CC	rgto	CGTGG
1 5 0	, ,	رى ت		112	GCC	CAI	יייטאני	, GG	F.	S	v	T	E	C	Q D	s	K	D	S	T
601	l Tr	ACA	GCC	TC	A GCA	GCA	CCCI	r GA	CGC	TGAG	C A	AAG	CAG	AC'	r ACG	AGA	AACA	CA	AAG'	TCTAC
		<b>700</b>	~~~	300	D CC4		CCCI	\	יכרכ	ልሮሞሮር	G T	TTC	GTC	.TG/	A TGC	TCT	1101	. 61	110	
	8 Y	S	L	. :	s s	r	L	Т	' L	S	K	. А	ע	) :	Y E		п	K	•	-
66	1 G	CCT	GCG	AA	G TCA	CCC	ATC	A GG	GCC	TGAG	C T	CGC	CCG	TC	A CAA	AGA	GCTT	CA	ACA TGT	GGGGA CCCCT
	C	GGA	CGC	TT	C AGI	GGG	TAG'	r cc	:CGG	ACTC	Aنی د	GCG P	<b>UU</b> U	.AGʻ	r Gr	S	F	N	R	CCCCT
72	1 6	ΔСΤ	ייים	מבי	G CTC	ATC	CTC	r ac	CGCC	GGAC	G C	ATC	GTG	GC	C CTA	GTA	CGC	A AC	TAG	TCGTA
, 2	_ G	TCA	CA	\TT	C GAG	TAC	GAG.	A TO	CGC	CCTG	C	TAG	CAC	CG	G GAT	CAT	'GCG'	r TG	ATC	AGCAT
21	8 E																			
		_								-1/			$\frown$	•						

FIG. 36

#### 45/136

	AAAAC	:CC	TAT ATA	C'	TAGA ATCI	GG'	TTG AAC	AGG	OTS CAC	TA:	AAA	TAC	TT.	${f r}{f r}{f r}$	CT	TATA ATAT I	GCG	TAA	AGA	AGAA	ICGI.
-1												11			••	-	•••	_	_	_	
841	TCTAT	GT	TCG	T	TTTI	TC'	TAT	TGC	TP	CA	AAC	GCC	TA(	CGC	TG	AGGT	TCA	GCT	AGT	GCAC	STCT
	ACS TE	(C)	ACC	Α		DA	ልጥል	ACC	:AI	rgt	TTG	CGC	YTA:	GCG	AC	TCCA	AGT	CGA	TCA	CGTC	JAGA
-11	S M	F	. v	•	F	S	I	A	7	ŗ	N	A	Y	A	E	V	Q	ъ	٧.	Q	3
001	GGCGG	-mc			ver	מים:	CCC	λCC	200	360	тса	CTC	CG	ттт	GT	ССТС	TGC	AGC	TTC	TGG	CTAC
	CCCCC	280	CCC	: A	CCAC	ىلىت	CGC.	TCC	$\mathbb{C}^{\mathbb{C}}$	CCG	AGT	GAC	3GC.	<b>AA</b> A	CA	GGAL	ALG	TCG	MAG	WCC,	SAIG
8	G G	G	3 L	,	V	Q	P	G	(	3	S	L	R	L	S	С	A	A	S	<u>G</u>	<u> </u>
061	TCCT	ויייו	ירכז		ያፈጋጥ	атс	тат	GC	AC:	rgg	GTC	CG'	rca	GGC	:cc	CGGG	TAA	.GGG	CCT	GGA.	ATGG
	ACCA		יככיז	, (	ነ አርጥ(	TAS	מידמי	CG'	TG	ACC	CAG	GC	AGT	CCG	GG:	GCCC	TTA:	CCC	GGA	CCI.	IACC
28	S F	- 5	55		Н_	<u>Y</u>	_M_	н	, 'i	M	V	R	Q	A	P	G	K	G	L	E	W
	GTTG	•																			
1021	GTTG(	GA?	TAT!	I A	T'TGA'	TCC	TTC	CA.	ATY Ta	CC 2	CTT	TG	I MC ÀŤG	CAT	יארי אין	TAG	ryri T	CAA	GTT	CCC	GGCA
4 Ω	V G	CTY	ATAT	[ } [	MC1.	nGC P	saag S	N	IA	G	E	T	_T_	<u> </u>	N	0	ĸ	F	K	_G	R
1081	TTCA	CT'	TTA	rc	CTCG	CGA	CAA	CT	CC	<b>AA</b>	LAAC	AC	AGC	ATA	ACC	TGC	AGAT	rgaa Cooo	CAG	CCT	GCGT
	AAGT	GA.	AAT	A (	GAGC	GCI	rgtt	GA	GG	TT"	TTTG	TG	TCG	TA'	rgg	ACG'	rcty M	YC.T.T.	GIC	.GGA T.	R
68	F T	1	L S	S	R	D	N	S		K	N	Т	Α	Y	L	Q	PI	14	3		••
11/1	GCTG	AC	GAC	<b>A</b> (	CTGC	CGT	гста	тт	'AC	TG:	rgca	AG	AGG	GG	ATT	ATC	GCT	ACAA	TGC	TGA	CTGG
	CCAC	TO	כיתכי	r (	SACG	CC	TADA	' AA	TG	AC	ACGT	TC	TCC	CC'	TAA	TAG	CGA'	$\mathbf{\Gamma}\mathbf{G}\mathbf{T}\mathbf{T}$	ACC	ACT	GACC
88	A E	:	D '	T	A	v	Y	¥	•	С	Α	R	G.	ם	Y	R	<u>Y</u>	N_	G_	D	W
	TTCT								~ A	200		СП	יראר	ירכ	יירית	ССТ	CGG	CCTC	CAC	CAA	.GGGC
				~	1010	-	~ > ~ ~	ነ ጥር	~~~	ישיים	こにかい	$\sim$ 2	CTC	こここ	AGA	GGA	GCC	GGAG	GIL	דו ביס	CCCG
108	F F	,	D	v.	W	G	Q	- 0	;	T	L	V	T	V	S	S	A	S	T	K	G
1261	CCAT	CG	GTC	T	TCCC		TGG	AC		TC	CTCC	A.P	IGA(	CA	CCI	GAC	CCC	CGTG	TC	GCCC	GGAC
100	GGTA	\GC	CAG	A	AGGC	.فاقاد . ۲	ACCI	יו כ ז	3 3	S	S	ĸ	S	T	5	G	G	T	A	A	L
1321	GGC	rgc	CTG	Ġ	TCA	\GG	ACT	A, C	rTC	ccc	CGAA	CC	:GG	TGA	CGC	TGI	CGT	GGAA	CT	CAG	SCGCC
	CCG	100	בתאר		ACTT	rcc	TGA'	r G	AA(	3GG	GCTI	' G(	GCC.	ACI	GCC	CACA	IGCA	CCTT	. GA	GTC( G	-0000
148	3 G (	2	L	V	K	D	Y	1	F	P	E	P	V	T	' '	, 5	; W	N	5	G	A
1381	ו כיייכו	שרנ	יאכר	c	GCG	rgc	ACA	כ כי	TT	ccc	GGCI	' G'	rcc	TAC	'AG	ר ככז	CAG	GACT	CT	ACTO	CCTC
	CAC	$r_{CC}$	2TCC	20	CGC	A C C	TOT	GG	AA	GGC	CCG	Y C	AGG	$\mathbf{A}\mathbf{T}\mathbf{C}$	TC	4. GG4	ICIC		4 GA	IGV	300A0
16	B L '	ľ	S	G	V	H	T		F	P	A	V	L	Ç	2 :	5 5	S G	L	Y	S	ь
4 4 4		<b>.</b> ~ (			mc a	~~	ייייכר	c c	ሞር	<b>ሮ</b> ልር	בר א כנ (	- ф	ጥርር	GCZ	ACC	C AG	ACCI	'ACA'	r cT	GCA	ACGTG
	TOC	TCC	20 A C	7	እርጥ	വാ	יארה	ദേദ	AG	GTC	CTC	3 A	ACC	CG:	ľGG	G TC	1667	1.1.0.1.1	n Ga	LOJ.	IGCMC
18	8 S	s	v	v	T	7	, P	_	s	S	S	L	G		r (	Q :	r 3	, I	C	N	V
150	1 AAT	CA	CAA	GC	CCA	GCA	₹¥C₽		AA TT	GG.	LCGA(	C A	ለሪዶ ጥርባ	ኒሌሌ( ነጥጥ	CVV 21.1.	C TC	GGG'	ATT1	G AA	CAC	ACAAA TGTTT
20	TTA 8 N	いて H	K K	ى p	GGT S	رى. 1	1 J	'	K	V	D.	K	K	( 7	V	E :	P I	ζ S	C	: D	K
156	1 ACT	CA	CAC	ΑT	GCC	CG	CCGI	'G C	:CC	AG	CACC	A G	AAC	TG	CTG	G GC	GGC	CGCA	T GA	AAC	AGCTA
	ma s			T 2	CCC		CCC	$\sim$	ഹ	יתרו	CTCC	тс	TTT	iac	GAC	C CG		こしじょ	W C		100112
22	8 T	H	T.	С	P	• ]	ħ (	•								<u>.</u>	·		•	- *	L
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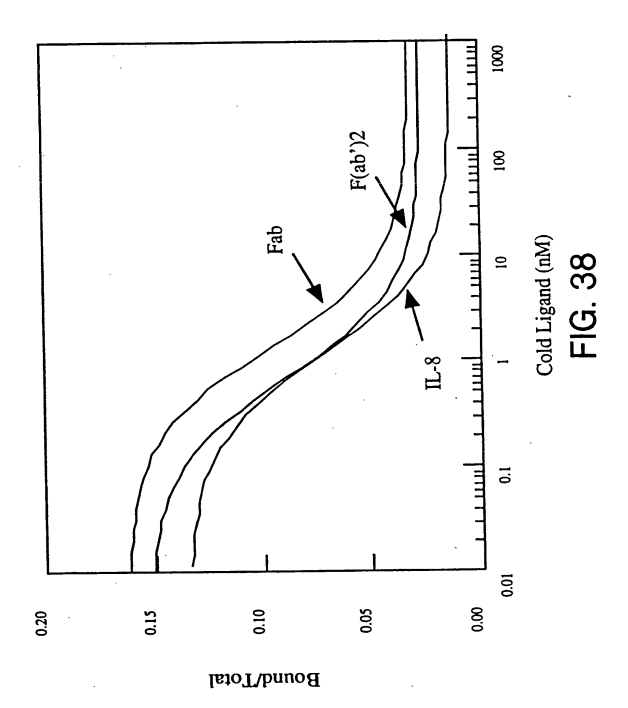
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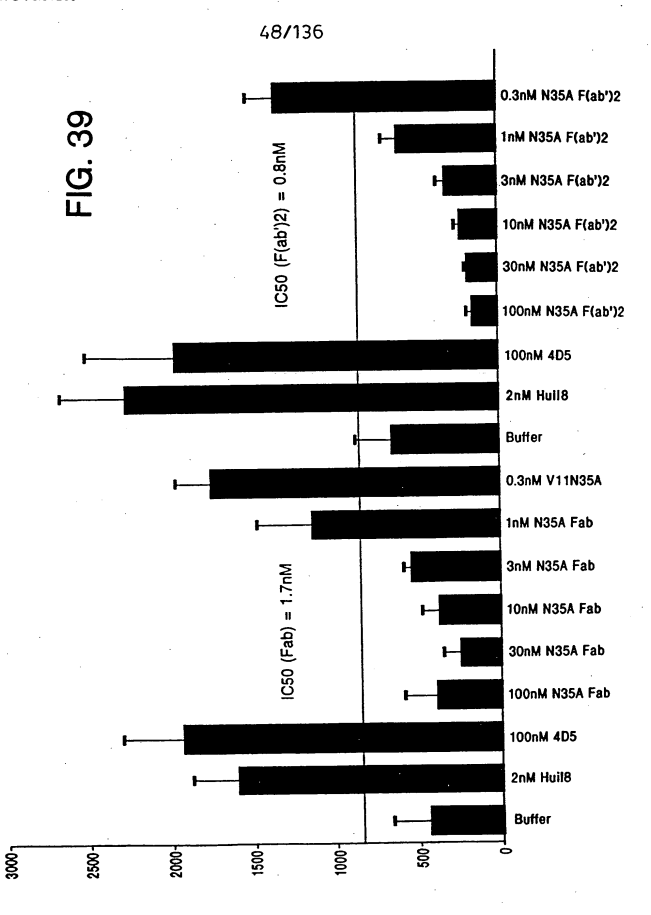
1621 GAGGACAAGG TCGAAGAGCT ACTCTCCAAG AACTACCACC TAGAGAATGA AGTGGCAAGA CTCCTGTTCC AGCTTCTCGA TGAGAGGTTC TTGATGGTGG ATCTCTTACT TCACCGTTCT 248 E D K V E E L L S K N Y H L E N E V A R

1681 CTCAAAAAGC TTGTCGGGGA GCGCTAA
GAGTTTTTCG AACAGCCCCT CGCGATT
268 L K K L V G E R O

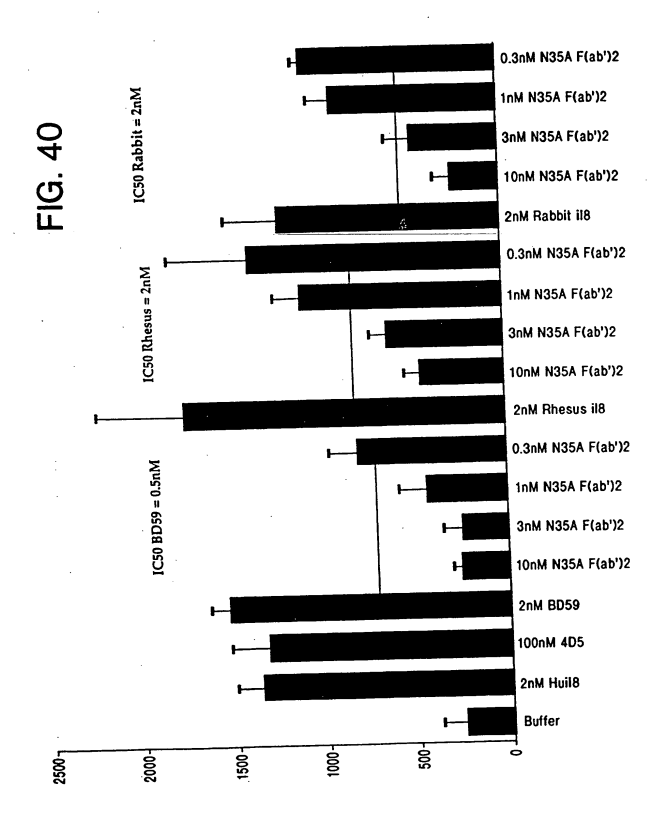
FIG. 37B



SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)

			I I DHI
alui prei hindili mboli taqi trugi eari/ksp6321 msel cac@l mboli hinfi GTTGTTATTT AAGCTTGCCC AAAAAGAAGA AGAGTCGAAT CAACAATAAA TTCGAACGG TTTTTCTTCT TCTCAGCTTA sau3Al	dpni(dam+) acii dpnii(dam-) nspBii bcli(dam-) mbli ACCAACAGGG GTTGATTGAT CAGGTAGAGG TGGTTGTCCC	mnli foki sfani TTGAAGCATC CTCGTCAGTA AACTTCGTAG GAGCAGTCAT	aiui ssti saci hgiJii hgiJii hgiJiii ecll36ii ecoRi bsp1286 rmai bsiHKAI maei bmyi bfai taqi maeIII apoi banII rTTGTAATTCGAGC
aluI hindIII 19I 11 cac8I AAGCTTGCCC A TTCGAACGGG I	acil nspBii ACCAACAGCG (		tru9I msel rtttaatgta
	hinPI hhal/cfoI g GGGAAATG c GGGTTTTAC	thal fnuDII/mvnI fnuHI bsoFI maeII bbvI bstUI snaBI bsoFI bsh1236I bbvI hinPI bsaAI aluI hhaI/cfoI GAGCTGCTGC GCGATTACGT AAAGAAGTTA CTCGACGACG CGCTAATGCA TTTCTTCAAT	IT TGTTTTATT AA ACAAAAATAA
ddeI bsrDI TCATIGCTGA AGTAACGACT	TCGCAATATG . AGCGTTATAC		pali III/eclXI ahdi/eamil05i imAi GACTT ATAGTCGC
nlaiii AAATACAGAC ATGAAAAATC TTTATGTCTG TACTTTTAG	msli maeili bsrDi ATTATCGTCA CTGCAATGCT TCGCAATATG TAATAGCAGT GACGTTACGA AGCGTTATAC	BI bsml A GCATTCCTGA CGACGATACG I CGTAAGGACT GCTGCTATGC	haeIII/ mcri eagl/xma eaeI cfri bsiEI naeIII bs
ecori pfimi apoi bsli GAATICAACT TCTCCATACT TTGGATAAGG CTTAAGTIGA AGAGGTATGA AACCTATTCC bspMi	hinPlhal/cfolhalulmstl alulavillyfspl hindill gaactgrgrg cgcaggraga agcTTGGAGCTCTCTCTCTCTTCGAGCTCTTCGAAACCTCT	rsal hinpl hbal/cfol mnll cac81 haell csp61 sfaNI GGGCGCTGTA GCAGGTATCC GGGCTACGGT	alui pvuli nepbii cttttcaaca getgteataa gaaaagtigt cgacagtatt
pflMI bslI r TCTCCATACT A AGAGGTATGA bspMI	hinPi hhal/cfol mstl avill/fspl l rg cgcaggragi	rsal hinpl hhal/cfol mnll haell csp61 gcgcccrgra CCGCGTAAAG	
ecori apoi 1 GAATTCAACT CTTAAGTTGA		<b>▼</b> =	tru9I mseI 301 KRAKGTIAAT TITICAATTA
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		SORPHINE SUFFI	···

FIG. 41A

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ddeI nlaIII
                                                                                                                                                                                                                                                                                                      501 ATACGCTGAT ATCCAGATGA CCCAGTCCCC GAGCTCCCTG TCCGCCTCTG TGGGCGATAG GGTCACCATC ACCTGCAGGT CAAGTCAAAG CTTAGTACAT
                                                                                                                                                                                                                                                                                                                        TAIGGGACTA TAGGICTACT GGGICAGGGG CICGAGGGAC AGGCGGAGAC ACCCGCTAIC CCAGIGGIAG IGGACGICCA GIICAGIIIC GAAICAIGIA
                TOGGIACCOS GGGAICCICI CGAGGIIGAG GIGAITITAI GAAAAGAAI AICGCAITIC IICIIGCAIC IAIGIICGII IIIICIAIIG CIACAAAGG
                                AGCCAIGGGC CCCTAGGAGA GCICCAACIC CACIAAAIA CIIIICITA IAGCGIAAG AAGAACGIAG AIACAAGCAA AAAAGAIAAC GAIGIIIGCG
                                                                                                                                                                                                                                                                     alui rsai
hindiii csp6I
                                                                        The penultimate nucleotide was changed fr {\tt G} tor {\tt ^{\circ}}
                                                                                                                                                                                               bspMI
                                                                                                                                                                                                                                                      sse8387I
                                                                                                                                                                                                                               pstI
                                                                                                                                                                                                                scfl
                                                                                                                                                                                                                                                                                             hphI bsgI
                                                                                                                                                                                                                                                                       DSpMI
          sfani
                                                                                                                                                                                                                                                                          maeIII
                                                                                                                                                                                                                                                        hphI
                                                                                                                                                                                                                                                                                             bstEII
          IIpoII
                                                                                   mutation was found that inactivated the mluI site.
                                                                   M
M
                                                                                                                                                                                  hgial/aspHI
                                                                                                                                                                                                    ecl136II
                                                                                                                                                                                                                     bsp1286
                                                                                                                                                                                                                                             bsinkAI
                                                                                                                                                                 hgiJII
                                                                                                                                                                                                                                                                                                     tth1111/aspI banII
                                                                                                                                                                                                                                                                                   bsrI aval aluI
                                                                                                                                                                                                                                                                  bmyI
                                                                                                                              sstI
                                                                                                                                                 saci
hphI
                                                                                                                                                                                                                                                                DSMFI
  bamHI aval
   asp718
                                           401
                                                                                 -23
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mbol/ndell[dam-]

sau3AI taqI

moli

bsaJI

avaI

cauli

xhoI

xmaI/pspAI

BCLFI

ncii dsav

smaI

hpaII dsav

Idsm

SCLFI

ncil

nlaIV paeR7I

dpn[[dam+]

csp6I rsal

nlaIV

kpnI cauII dpnII[dam-]

bstYI/xhoII

hgiCI

bani bsaJi alwi[dam-]

	52/136	
tfii hinfi bsmFI clal/bsp106 pleI bspD1[dam-] hinfi crgatttaca agtatcca rcgatrcrcr gaggrcccrr gactaaargr ircataggr agctaagaa L I Y K V S N R F S G V P S	rsal csp61 scal nlaIII CGCAACTTAT TACTGTTCAC AGAGTACTCA GCGTTGAATA ATGACAAGTG TCTCATGAGT A T Y C S Q S T H	acil mboli ATCTTCCCGC CATCTGATGA GCAGTTGAAA TAGAAGGGCG GTAGACTTT I F P P S D E Q L K
CIGATITACA GACTAAATGI L I Y K	mboli bpuAi bbsi CAGAAGACTT GTCTTGAA E D F	mboli bpuAl bbsI ATCTGTCTTC TAGACAGAAG S V F
TCCGAAACTA AGGCTTTGAT P K L	fnu4HI bsoFI bbvI scfI pstI bsgI AGTCTGCAGC TCAGACGTCG	] fnu4HI bsoFI bbvI ; TGGCTGCACC
scrfi mval ecoRII dsaV bstNI aluI apyI[dcm+] CAACAGAAAGC TCCGAAACTA GTTGTCTTG GTCCTTTTCG AGGCTTTGAT	TCT GACCATCAGC	sau3AI mbol/ndeII[dam-] fnu4HI dpnI[dam+] bsoFI dpnII[dam-] bbvI GGTGGAGATC AAACGAACTG TGGCTGCACC CCACCTCTAG TTTGCTTGAC ACCGACGTGG V E I K R T V A A P
	G ATTCACTCT	
bsri ACACTGGTAT TGTGACCATA E W Y	II[dam-] +] m-] -] -] bsmFI   rcrGGGACG	styl bsaJI rsaI csp6I csp6I nlaIV kpnI hgiCI panI asp718 acc6SI acc7IIGGAC AGGTACCAA IGCAAACCIG ICCCAIGGIT I F G Q G I K
CTACGTATTT GATGCATAAA T Y L	mspI hpalI bslI bsaWI sau3AI mbol/ndelI[dam-] dpn[[dam+] dpnI[dam-] alwI[dam-] alwI[dam-] nlalV bstYl/xholI bamHI alwI[dam-] bamHI alwI[dam-] cGGATCCGGT TCTGGGACGG ACCTAGGCCA AGACCTGCC G S G T D	maell ACGTTTGGA TGCAAACCT
GGTATAGGTG CTACGTATTT CCATATCCAC GATGCATAAA G I G A T Y L	CTCGCTTCTC GAGCGAAGAG R F S	bsrBI acii bsmFi TGTCCCGCTC ACAGGGGAG
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GAIGITIGG CATGCGACTC CAAGICGAIC ACGICAGACC GCCACCGGAC CACGICGGIC CCCCGAGIGA GGCAAACAGG ACACGICGAA GACCGAIGAG
                                                                                                                                                                                                                                           1301 CINCANACGO GIACGOTGAG GITCAGOTAG IGCAGIOTGG CGGIGGCOIG GIGCAGOCAG GGGGCICACI CCGITIGICO IGIGCAGOII CIGGCIACIO
                                1201 AGTACGCAAC TAGTCGTAAA AAGGTATCT AGAGGTTGAG GTGAITITAT GAAAAAGAAT ATCGCATITC TTCTTGCATC TATGTTCGTT TITTCTAITG
                                                                                                                                                                                            alwNI[dcm-]
                                                                                                                                                                                                          fpu4HI
                                                                                                                                                                                                                      bsoFI
                                                                                                                                                                                                                                    bbvI
                                                                                                                                                                                    aluI
                                                  TCAIGCGIIG AICAGCAIII IICCCAIAGA ICICCAACIC CACIAAAAIA CIIIIICIIA IAGCGIAAAG AAGAACGIAG
                             sfani
                            Ilodm
                                                                                                                                                                                                bsp1286
                                                                                                                                                                                                                                         banII
                                                                                                                                                                                                              apyI[dcm+] bsaJI bmyI
                                                                                                                                                                                                                          haeIII/pall apyI[dcm+]
                                                                                                                     ecoRII
                                                                                                                                                                                     dsav bstNI hgiJII
                                                                                              scrFI
                                                                                                                                   dsaV
                                                                                                          mvaI
                                                                                                                                                            fnu4BI
                                                                                                                                                                                                   bstNI bsoFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ecoRII
                                                                                                                                                                                                                                                                                                                                                                                                                                                       SCIFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   mvaI
                                                                                                                                                                          ecoRII
                                                                                                                                                  SCIFI
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rmaI
            maeI
                        bfaI
                                                                                                                                                                                                                      maeI
                                                                                                                                                                                                                                  bfaI
                                                                                                                                                                                                           rmaI
                                                                                                                                                                                                                                                                                     .a
                                                                                                                                                                                                                                   mluI csp6I mnlI
                                                                                                                                                                      bsiWI/splI
                                                                                                                                                                                               EnuDII/mvnI
                                                                                                                                                           rsal
                                                                                                                                                                                                                         bsh1236I
                maeI
                            bfaI
                                                                                                                                                                                                            bstol
     rmaI
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haeIII/pall eco01091/draII

nlaIV apyI[dcm+] bsmAI

bsaJI hphI bsmBI

moli

esp3I

bstNI

ahaII/bsaHI

styl asul bsaJI TCGGCCTCCA CCAAGGGCCC

	33713	O	
maeII I bsaAI C TACGTATAAT G ATGCATATTA T Y N	T GCCGTCTAIT A CGGCAGAIAA A V Y Y	sau96I haeIII/palI sau96I nlaIV hgiJII bsp1286	bmyI banII asuI apaI
m-] snaBI m-] hphI CCA ATGTGAAAC GGT TACCACTTTG N G E T	cac81 mnli I ddel drdi GCGTGC TGAGGACAC CGCACG ACTCCTGTG		moli [
bsli sau3Al mbol/ndell[dam-] dpnl[dam-] dpnll[dam-] ATATT GATCCTTCCA TATAA CTAGGAAGGT I D P S N	cacé cacBI GAACA GCCTGCG CTTGT CGGACGCJ N S L R	maeIII	scrii scrii mvai ecoRII bsaJI dsav bseRI
I GGGT TGGAT CCCA ACCTA V G Y	fnuDII/mvnI scfl bstUI cac8I mnlI bsh1236I bspMI cac8I ddel drdI bsh236I bspMI cac8I ddel drdI uI cgcgacaact ccaaaaacac accatactg cagatgaaca gcctgcgtgc tgaggacat gccgtctatt gcgctgttga ggtttttgtg tcgtatggac gtctacttgt cggacgcaca actcttgta cggcagataa gcgctgttga ggtttttgtg tcgtatgac gtctacttgt cggacgcaca actcttgta cggcagataa		
bsaJI dsaV avaI bstNI bsaJI bslI sau961 apyI[dcm+] nlaIV sau961 haeIII/palI asuI eco01091/draII haeIII/palI AGGCCCG GGTAAGGGC TGGAATGG TCCGGGG CCATTCCCG ACCTTACC	CCAAAAACAC AG GGTTTTGTG TC K N T A		maell hinll/acyl
	T T		
sau96I avali asul blaly bsrl iGC ACIGGGCCG	haeIII/palI au96I suI iGGCCGTTT CACTITATC iCCGCCAAA GTGAAATAG		
pleI hinfI sau96I taqI avaII xhoI paeR7I bsrI avaI maeIII bsrI GAAGAGCTCA GTGATATACG TGACCCAGGC	haeIII/palí sau96I asul 1501 CAAAAGTTCA AGGGCGGTT CACTTTATCT GTTTTCAAGT TCCCGGCAAA GTGAAATAGA		
taq taq xhoI paeR avaI 1401 CTTCTCG GAAGAGC	1501 CAAAAG GITITG		

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TGACACGITC ICCCCIAAIA GCGAIGITAC CACIGACCAA GAAGCIGCAG ACCCCAGITC CIIGGGACCA GIGGCAGAGG AGCCGGAGGI GGIICCGGG C A R G D I R I N G D W F F D V W G Q G I L V I V S S A S I R G P

1601 ACTGTGCAAG AGGGGATTAT CGCTACAATG GTGACTGGTT CTTCGACGTC TGGGGTCAAG GAACCCTGGT CACCGTCTCC

mboll aatil tagI

hphI bsrI maeIII

moli

sed right is from p6G425chim2.fab2

	56 /136	suI[dcm-]	
hphi nspi hpali cfr101/bsri bsawi tth bsli agel maeli TCCCCGAACC GGTGAC	> H > A B B B B B B B B B B B B B B B B B B B	ddel fnu4HI maeIII mnlI bsoFI maeIII mnlI bsoFI maeIII mnlI ddel hphl bsp1286 cc0811 hinf! ddel bbvl bstEII bmyl bpml/gsul[dcm-] cctacagtcc tcacactct actcctcag cagcgragtg accgracct accctcag cagcarcac ragcacacac ragcacacacac ragcacacacacacacacacacacacacacacacacacac	hglJII bsp1286 Lidii bmyi maelii Danii maelii GAAAGTIGAG CCCAAATCTT GTGACAAAAC CTTTCAACTC GGGTTTAGAA CACTGTTTTG
scrFI mval scrFI mval ecoRII dsaV bstNI ecoNI bstNI ecoNI bstNI fnu4HI bsoFI bsaJI bslI acil apyl[dcm+] bsp1286 asuI bsoFI bmyl nspBII bsaJI bbvI apyl[dcm gcgcGCACAG CGGCCCTGGG CTGCCTGGTC ccccGTGTC GCCGGACCC GACGGACCAG	GGTAALGCLV	mspi hpali scrfi ncii dsav cauli rCCCGCTGT AGGCCCACA	taqi sali fi styi hincii/hi Ei bsaji acci rcacaacacca aggrcgacaa agrgrrcgg rcgrrgrgr E R P S N I R V D R
	GGGGCCGTG GGAGGAGGII P L A P S S K hinPI hhal/cfol	hari kasi cac hinli/acyi cac hgiCi fnu4Hi haeli bsoFi bani acii el ahali/bsaHi nspBli cAGGCGCCT GACCAGCGC GTCGCGGGA CTGGTCGCCG	
	TAGCCAGAAG 129 S V F	dde 1801 TCGTGGAACT AGCACCTTGA 162 S W N S	aluI fnu4HI bsoFI bbvI bstXI 1901 CCAGCAGCTT GGTCGTCGAA

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FIG. 41G

fnu4HI bsoFI  mcrI  mcrI  eagl/xmalIl/eclXI  eagl/xmalIl/eclXI  eagl/xmalIl/eclXI  eagl/xmalIl/eclXI  bsiEI  nbelI  nbelI  nspl  nsp	scrFI  I mael hpall hpall sau961 pleI thu4BI haelII/palI bsoFI asuI hinfI bsoFI cGACGCCCT AGAGTCCTA ACGCTCGGTT GCCGCCCC CAAAAAATAA	tru9I msel taqI hindIII hpaI nlaIII claI/Dsp106 tru9I hincII/hindII aluI bspDI[dam-] msel aciI hincII/hindII aluI bspDI[dam-] msel aciGGTAGTT TATCACAGTT AAATTGCTAA CGCAGTCAGG CACCGTGTAT GAAATCTAAC GTTAACTCAT GTTTGACAGC TATCATAGTTAAA ATAGTGTCAA TTTAACGATT GCGTCAGTCC GTGGCACATA CTTTAGATTG
fnu4EI bsoFI haeII/palI mcrI eagl/xmaIII/eclXI eagl/xmaIII/eclXI eaeI cfrI bsiEI notI bsoFI bsoFI naeI bsoFI acit aciI acit aciI acit aciI acit aciI L G G R M R Q L E D R y and leucine zipper	sphi ddei nlaili celli/espi rmai blpi/bpull021 maei hinPi nspi bfai bsmFi hhai/cfoi sau96i plei haeii nspHi haeiii/pali i eco47111 cac81 asui hinfi crcgcgcacc gctaaccatc ccacccta v G E R O	tru9I  msel  taqI hindili  hpal nlaliI  clal/bspl06 tru9I  hincil/hindiI alui bspDI[dam-] msel acil  msel  hincil/hindiI alui bspDI[dam-] msel  GTIAACTCAT GTTTGACAGC TATCACAGT AAGTGTAA TATCACAGTT AAATTGCTAA CGCAGTCAGG CACCGTGTAT  GATTAACTCAT GTTTGACAGC TAACTAGCT ATCGAAATT ACGCCATCAA ATAGTGTCAA TTAACGATT GCGTCAGTCC GTGGCACATA
fnu4HI bsoFI haell/pa mcri eagl/xmalli eagl/xmalli eagl/xmalli eagl/xmalli eagl/xmalli eagl/xmalli eagl/xmalli eagl/xmalli spri noti fnu4HI noti noti fnu4HI bsoFI noti acit bmyl acit bmyl acit bmyl scoccarcc caccacarc caccacarc acit acit hal bsoFI cacacarcc caccacarcc caccacarc acit acit acit bmyl acit bmyl acit acit bmyl acit acit fnu4HI bsoFI bsoFI bsoFI cacacarcc caccacarcc caccacarc acit acit cacacarcc caccacarcc caccacarc acit acit acit acit cacacarcc caccacarcc caccacarc acit acit acit acit acit acit bmyl acit	sphi ddei nlaii celli/espi blpi/bpull0 hinPi nspi hhal/cfol hinfi hindiii eco47111 cac81 ctcttactic accgifcaa Gitticaa Gitticaaci	tru91 msel hpal nlaIII aluI bspD. 2201 GTTAACTCAT GTTTGACAGC TTATCATGGC CAATGAGTA CAAACTGTGG AATAGTAGC

ecoRV ecil cttgcgggar atcgtccaft gaacgccta tagcaggtaa	fnu4HI CO bsoFI bsoFI > 1286 HKAI mcrI eael C I bsiEI cfrI O GCACTGTCG ACGGCTTTGG	sau3AI mboI/ndeII[dam-] dpnI[dam+] dphI[dam-] alwI[dam-] llaIV sstYI/xhoII hgaI samHI mspI llwI[dam-] hpaII sfaNI iWI[dam-] hpaII sfaNI iGA TCCTCTACGC CGGACGCATC
haelli, sau961 ncil rsal mspl mnll csp61 hpall hpall caull cfr101/bsrFl asuI ATGCCGTAC TGCCGGCCTA	hinp!  hhal/cfo!  rma!  mae!  nhe!  4HI hae!I  Hinp!  hhal/cfo!  bsp1286  acil hae  4HI hae!I  hhal/cfo!  bsp1286  cacl!  r bfa!  avill/fsp!  bmy!  bsiE!  cft:  cacl!  rGCTAGCGCT ATATGCGTTG ATGCAATTTC TATGCGCACC CGTTCTCGGA GCACTGTCCG ACCGCTTTGG  ACGATCGCGA TATACCAACTTAAAG ATACGCGTGG GCAAGAGCCT CGTGACAGC TGGCGAAACC  mni!	.idam+] .[dam+] .[dam-] .ni .ni .ni .ndGC GACCACACC GTCCTGTC .nacc CTCTGTC .nacc CTCTCTC .nacc CTCTCTC .nacc CTCTCTC .nacc CTCTCTC .nacc CTCTCTCTC .nacc CTCTCTCTC .nacc CTCTCTCTC .nacc CTCTCTCTC .nacc CTCTCTCTC .nacc CTCTCTCTC .nacc CTCTCTCTCTC .nacc CTCTCTCTCTC .nacc CTCTCTCTCTCTC .nacc CTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTC
sfaNI scrFI mvaI ecoRII dsaV bstNI bsaJI maeIII fokI scfI cGTC ACCCTGGATG CTGTAGGCTT GCAG TGGGACCTAC GACATCCGTA TCCGAACCAA	hinPI hhal/cfol rmal mael nhel fuu4HI haeII bsoFI eco47III bbvI bfaI SfaNI ic cac8I stc TGCTAGCGCT ATATGCGTTG ATGCT	nlaIV TGGAGCCACT ACCTCGGTGA
sfaNI scrFI mvaI ecoRII dsav nlaIV bstNI mnlI hglCI bsaJI hhaI/cfoI fokI banI maeIII fokI scfI hhaI/cfoI cGCACCGTC ACCTGGATG CTGTAGGCAT TTACGCGAGT AGCAGTAGGA GCCGTGGAGGAT	hinPI hhal/ci rmal hhal/ci rmal mael nheI nheI sfaNI bsrI sfaNI bsrI cac8I cac8I cac8I gGCTGTCGTA GCGTCAGTGCTAGCGCT GGCTGTCGTAGCGGAGGATCGCGA	acil fnu4Hi bsoFi acil bsri cac8I 2501 CCGCGCCCA GICCTGCTACT GGCGGCGGGT CAGGACGAGC GAAGCGATGA
	SUBSTITUTE SHEET	

bsrI alul bslI

sfaNI

hgaI

pleI hinfi

ecoNI bslI

bsrI bbvI

2801

	59/136	
II cfoI	I haeIII/palI GG CC	
rcal hinpi hgiJii haeli bsp1286 eco47III bmyl bspHl hhal/cfoI cgc crcArGAGCG	aspRI 36 AI LCAACGG AGIIGCC	hpail bsawi
rcal h hgijii ha bsp1286 ec bmyl bspHl h banii nlaiii iGGG CTCATGAG	fnu4HI bsoFI hgiAI/aspHI acii bsp1286 fnu4HI bsiHKAI bsoFI bmyI acii acii ha GCGGCGGGG TGCTCAACGG	d Thert alut half
am-] CCACTTC	fnu4HI bsofi acii fnu4HI bsofi acii aci gcGGCGCG	t r
hgiJII bsp1286 bmyI banII sau3AI cac8I mboI/ndeII[dam-] hgiJII haeII dpnI[dam+] bsp1286 eco4' dpnI[dam-] bmyI bspRI hhai mboII[dam-] banII nlaIII GATGGGGAAG ATCGCGCTCG CCACTTCGGG CTCATGAGCG	bsli accaticcit tggtaaggaa	
sa mp dt mboll sarggggaag	cac81 CCTTGCACGC	FX 64
	hinPI hhal/cfoI nlaIV narI kasI hinlI/acyI haeII ahaeII GGGGCCATCI CCCGGGTAGA GGAACGTGGG	•
hinPI hhal/cfoI nlaIV narI kasI hinlI/acyI hgiCI aalI/bsaHI eBI	Pall banfi GGACTGTTG	
hinpi hhal/cfol nlaly nari kasi kasi hinli/acyi hgiCi haeli bani ahall/bsaH cac81	scrFI scrCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	pleI
II acil	dsa gau961 nla1v hae111/k asu1 bst eco01091/ cac81 bs11 GCAGGCC CC	ecoNI
hhal/cfol hhal/cfol hhal/cfol hall hall hall hall hall hall hall ha	CACCGGCCGT AGTGGCCGCG GTGTCCACGC CAACAACGC GGATTTTCGG GTGTCCACGC CAACAACGC GGATTTTCGG GTGTCCACGC CAACAACGCC GGATTTTCGG CGTGGCCGG GGCACCTGCGGGCCG GGGACTGTTG GGAGCTGTGG GGACTGTTG GGAACGTGCGG GGCACCGGCCC CCTGACAACC CCGCGGTAGA GGAACGTGCGGGCCGCGCGCGCGCGCGCGCGCGCGCGCGC	fnu4HI bsoFI
	CACCGGCCGT .	
2601	270	

CCTCAACCTA CTACTGGGCT GCTTCCTAAT GCAGGAGTCG CATAAGGGAG AGCGTCGTCC GATGCCCTTG AGAGCCTTCA ACCCAGTCAG CTCCTTCCGG GGAGTTGGAT GATGACCCGA CGAAGGATTA CGTCCTCAGC GTATTCCCTC TCGCAGGAGG CTACGGGAAC TCTCGGAAGT TGGGTCAGTC GAGGAAGGCC

acil thai thai thai thai thai thai thai thai	sau961 nlaiv nlaiv avaII avaII asuI cacl tfiI acacl mnlI maeIII bsmFI GCTTGCGGTA TTCGGAATCT TGCACGCCCT CGCTCAAGCC TTCGTCACTG GTCCCGCCAC CGAACGCCAT AAGCCTTAGA ACGTGCGGGA GCGAGTTCGG AAGCAGTGAC CAGGGCGGTG	mcri eagl/xmalil/eclXi eael hinPI cfri hhal/cfoi hgai nael fnu4HI fnuDII/mvnI cfri01/bsrFi bstUi cac8I acil hgal bg1I nlaIII haelII/palI cGCGGCGTAG GCGCGGCGGCGGCGGCGGCGGCGGCGGCGGCGGCGGCG
mboli bpual bbsi ttatgactg tcttctttat Catgcaactc Gtag	/pa dam dam trc	mcrI eagl/xmaill/eclXi eael hinPI cfrI hhal/cfoI nael fnu4HI fnuDII/mvnI cfr101/bsrFI bstUI I hpail bsoFI bsh1236I cac8I acii hgaI b91I nlaili haelII/palI maelI cGCCGCCATG GCGCCGAC GCTGGCTA CGTC
acil thai thai fuuDII/mvol bstul nlallI acil bsh12361 fou4HI hhai/cfol 2901 TGGGCGCGG GCATGACTAT CGTCGCCGCA C3 ACCCGCCCC CGTACTGATA GCACGGCGT G3	thai fouDII/mvoI bstUI haeIII bstUI haeIII sau96I hinPI mboI/ndeII[ avaII bpmI/gsuI[dcm-] dpnI[dam-] asuI bpmI/gsuI[dcm-] dpnI[dam-]	haeIII/palI haeI psp14061 cac81 3101 CAAACGTTTC GGCGAGAAGC AGGCCATTAT C

FIG. 41K

	617136	
alwI[dam-] G	•	
. G		•
. [A]	nlaIII CATG GTAC	ဗ္ ပ္
PAAG	CAT	mbli V V V V C C C C C C C C C C C C C C C
I TTC	1 1 1 1 1 1 1 1	14HI OFI II mu nlaIV hgiCI DanI CGGCACC
alu AGC	nlal GGIIGGCAIG CCAACCGIAC	fnu4HI bsori acii I nlaIV hgici OI/bsrFi I bani G CGGCACCTCG
bsmFI aluI GGGA CAGCT CCCT GTCGA		fnu41 bsoF. acii nspi hpali n. tael h. str101/b. str301/b.
# # # # # # # # # # # # # # # # # # #	/aspHI 86 AI nlaIII ACATGGAACG	fnu4HI bsoFI acil nspI m hpaII nlaIV naeI hgiCI cfr101/bsrFI cac8I banI GCGG CGCCAC
TCA	aspel 6 .i nlaiii cargga	GAA
bsmFI aluI a. CCATCAGGGA CAGCTTCAAG	(/as 1866 (Al 1017) (Al	TG
	hgial/aspHI bsp1286 bsiHKAI bmyi ncBi nlaII GGGC ACATGG	GAA
GAC	hgir bspl bsli bmyl cac8I	CCT
GAT	r TCG ,	alI taqI .I TCGA(
	muli bsaJi ii iii iii iii ii cro	haeIII/pall for sau961 scrfl ncil mspl hpall dsav taql cf1101 ccull mnll cac81 CCGGCCACC TCGACCTGAA TGGAAGCCGG
bspMI scrFI mvaI ecoRII dsaV bstNI apyI[dcm+] CCAGGCAGG	mn bs. acil fnu4HI bsoFI bg1I	fill fill cac
bs. scrfi mval ecoRII dsav bstNI apyI[d	b b TATG	haell sau961 crfl cil spl pall sav asul aull
bspMI scrFI mvaI ecoRII dsaV bstNI apyI[dcm+] TCCAGGCAGG TAGATGACGA AGGTCCGTCC ATCTACTGCT	mnli bsaJI hgiAI acii bsp12 fnu4HI bsiHK bsoFI bmyI bglI cacBI TTTATGCCGC CTCGGCGAGC	hae sau9 scrFI ncii mspi hpaii dsaV V asuI CCGGGC
ni haelli/pali 31 nlaili GGCCATGCTG 7 CCGGTACGAC 8	а 20. Ст. А 20.	Inlary Correction of the corre
/mvnl hael (61 haelli/pa cac81 nlalil cca ggccArgcr (cgr ccggtAcGA	sau3AI mbol/ndeII[dam-] I maeIII dpnI[dam+] dpnI[dam-] GATC GTCACGGCGA	vnI nlalI scATGGAC
elli CCA	I ndeII[c maeIII [dam+] [[dam-] GTCACG	/mvi 61 n TGCG
ivbI haeI haeI haeI 128 GG	61  I sau3AI mbol/ndeII[ dam-] nspBII maeIII acil dpnI[dam+] ccGCTGATC GTCACG	thai fnuDII/mvnI bstUI racii nla cGGG GTGCAT
thai fnubli/mvni bstUl hae 13 bsh12361 h acii cac81 CGGGTTGCA G	sau3AI mbol/n I m dpni[d dpni[GCATC G	thai fnuDi bstUI bsh12 ai acii GTCGG
II IDII IUI II 23	961 II Sa I mb [dam-] nspBII acii di ACGGCGGC	iva igaar Sogaar
thaI fnuDII/mvnI bstUI haeI bsh1236I haeIII/pal] aciI cac8I nlaIII CCGCGTTGCA GGCCATGCTG	sau96I avall bsrI sau3Al asuI nbol/ndeII[dam-] ipnI[dam+] nspBi iqnI[dam-] aciI sATCACTGG ACCGCI	haeIII/pall sau961 sau961 bs crFI thaI ncil mspl fnuDII/mvnI mspl fnuDII/mvnI mspl fnuDII/mvnI mspl hpaII hpaII bstUI bstUI dsaV bstUI bsh1236I nlaIV asuI taqI cfr101 mnlI acil hgaI acil nlaIII cauII mnlI cac8I ccrcccGcG TTGCGTCGC GTGCATGGAG CCGGCCACC TCGACCTGAA TGGAAGCCGG GGAGGGCGC AACGCAGCGC CACGTACCTC GCCCCGTGG AGCTGGACTT ACCTTCGGCC
cac81 Fr NI b GC CC	sau96I avali :I asul leII[da lm+] ns lam-] IGG ACC	thal fnuDII bstUI bshl23 cdI
ca sfaNI fokI GGGATGC	ss av bsri Al as (dam- [[dam-] lm-]	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC
cac sfani foki CGGGATGC GCCCTACG	sau9 avaI bsri bsri mbol/ndell[dam+] dpnl[dam+] taql[dam-] t GGATCACTGG A	n I I CTO GAĞ
14HI PFI I mslI sfani SGC ATC	S E O O CO	
fnu4HI bsofi acii ni ms iii sfa	t IGA	AGA
fou bsc aci mspI hpaII CCGGCC	ATE	TIG
fou4HI bsoFI acil mspl mslI hpaII sfaNI TTCCGCGCGC AT	sau96I avall bsrl sau3Al asul mbol/ndell[dam-] dpn1[dam+] nspBlI maeIII dpn1[dam+] acil dpn1[dam+] taq1[dam-] acil dpn1[dam-] TCGATTGAA GCTAGTGACC TGGCGACTAG CAGTGCCTT	ACCTIGICIG
ប្តូម	######################################	AT.
SAGO	4HI I. I. II/mvnI il idam-] GGCTCTTACC CCGAGAATGG	T I III I I I I I I I I I I I I I I I I
mboli : :: :TCTTC	fnu4HI bsoFI acil thai fnuDII/mvnI bstUI Bsh1236I erI[dam-] m+] cGC GGCTCTT GGC CCGAGAA	fnu4HI bsoFI hlnPi hhal/cfoI nlaIV narI kasI hinlI/acyI hgiCI banI aciI ahaII/bsaHI GGC GCGGGG
mt tfii hibfi GA TT		for bsc hlopi hhal/c nlary nari kasi hinli/i hgici haeri aali/l
tfii hinf ACT A	fpu4E bsori acil thai fnuDI; bstUI bsh12; leII[dam+] lam+]	fi bibDP hinP, hhal, nari kasi hinli, hgici haeli abali agge G
TTA	fou bso aci thal foul bstu cac8! cac8! sau3A! bshl abo!/nde!![dpn!] dpn![dam+] dpn![dam+] dpn![dam+]	fnu4BI bsoFI hinPI hhal/cfoI nlaIV narI kasI hinlI/acyI hgiCI haeII banI aciI ahaII/bsaBI CTAACATCCG CGGCGGATA
thai scrfi bsoFI fnu4HI fnuDII/mvnI mvaI bsoFI acil cac8I hael dsaV tfil mspI mslI sfaNI bsh1236I haelII/pall bstNI hinfI hpaII sfaNI fokl acil cac8I nlaIII apyI[dcm+] CCCATTATGA TTCTTCTCGC TTCCGGCGC ATCGGGATGC CCGCTTCCA GGCCATGCTC TCCAGGCAGG TAGATGACG GGGTAATACT AAGAAGACG AAGGCCGCTACG GGCGCAACGT CCGGTACGAC AGGTCCGTCC ATCTACGT	fnu4HI bsoFI acil thal thal fnuDII/mv bstUI cac8I sau3AI bsh1236I mboI/ndeII[dam-] dpnI[dam+] dpnI[dam+] dpnI[dam+] cTAGCGAGCG CGCM	fnu4HI bsoFI hinPI hinPI hhal/cfoI nlaIV narI kasI hinlI/acyI hgiCI haeII banI aciI ahaII/bsaHI CTAACATCCG CGCGGGATA TGGAACAGAC
mboli tfil hibfi 3201 CCCATTATGA ITCTTCGC GGGTAATACT AAGAAGAGCG		3401
32(	3301	34

hgal thal acil fnuDII/mvnI bstUl bshl2361 TCGCGTCCGC	cac8I  (/draII  aciI  aciI  cGATCCGACC
hinpl hhal/ofol hhal/ofol mstI pflMI avill/fspl styl bstUl binfl bsll nlaIV acil bsml bsll bsaJI bshl2361 3501 CTAACGGATT CACCACTCCA AGAATTGGAG CCAATCAATT CTTGCGGAGA ACTGTGGATG GCGAACCAA GCACTGGCAG AACATATCCA TGGCGTCGCG GATTGCCTAA GTGGTGAGGT TCTTAACCTC GGTTAGTTAA GAACGCTCT TGACACTTAC GCGTTTGGTT GGGAACCGTC TTGTATAGGT AGCGCAGGCG	msp hpall scrfl scrfl ncil dsav avall avall puMI mael ecool091 caull-bfal
cfol pflMI spI styI bslI bsaJI AAACCAA CCCTTGC	sau3AI mbol/ndeII[dam-] dpuI[dam+] dpuI[dam+] cfol hgiAI/aspHI laIII bspl286 aviII/fspl bsiHKAI nslI bmyl mn
hinPI hhal/cfol mstI pf avill/fspI bsmI bs ACTGTGAATG GGGAAACG	haeII/pall  mscl/ball haeI scrFI mval dsal ecoRII dsav bstNI bslI bsaJI mbol/ndeI apyI[dcm+] avalI hinPl dpnI[dam+] avalI mstl nlalII bspl286 nlary cfrI aviII/fspl bsiHK ecol091/draII mslI bmyI GGGTCCTGGC CACGGTGCG CATGATGTG C
acii CTTGCGGAGA GAACGCCICI	haeIII/I mscI/ball haeI scrFI mvaI dsaI ecoRII dsaV bstNI bstNI bstNI bslI bsaJI avaII avaII nlaIV cfrI ecol109I/draII gGGTCCTGGC CAC
/ CCAATCAATT GGTTAGTTAA	fnu4HI I bsoFI bbvI ; GGGCAGCGTI
II DlaIV AGAATIGGAG	fnu4HI thai hinPI bsoFI fnuDII/mvnI bstUI 8I hhai/cfoI acil sfaNI cGC GGCGCATCTC G
hphi pfimi tfii pfimi hinfi bsli carr caccactcca ac	fnu4HI thal hinPI thal hinPI fnu4HI bsoFI bsoFI fnuDII/mvnI fnu4HI bstUI bsoFI cac8I hhal/cf bbvI aciI bsh1236I a' bpmI/gsuI[dcm-] aciI sfaNI CTCCAGC AGCCGCACG GGGGCATG
tf bil 3501 CTAACGGA GAITGCCT	bpmI, GIAGAGGI

3701 CGGGGTTGCC TTACTGGTTA GCAGAATGAA TCACCGATAC GCGAGCGAAC GTGAAGCGAC TGCTGCTA AAACGTCTGC GACCTGAGCA ACAACATGAA GCCCCAACGG AATGACCAAT CGTCTTACTT AGTGGCTATG CGCTCGCTTG CACTTCGCTG ACGACGACGT TTTGCAGACG CTGGACTCGT TGTTGTACTT fnu4HI bsoFI bbvI fou4BI bsoFI bbvI bsh1236I maeII fauDII/mvaI bstul

cac8I

hphI

hinfi tfir

maell

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GACACCITGI GGAIGIAGAC ATAATIGCII CGCGACCGIA ACIGGGACIC ACINAAAAGA GACCAGGGCG GCGIAGGIAI GGCGGICAAC AAAIGGGAGI
                                                                                                                                                                                                                                                                                                                                        CCGCCAGIIG TITACCCICA
                                                                                                                                                                                                            ACCAGAAGCC AAAGGCACAA AGCATITCAG ACCIIIGCGC CIICAGICGC GGGACGIGGI AAIACAAGGC CIAGACGIAG CGICCIACGA CGACCGAIGG
                                                                                                                                                                                             IGGICIICGG IIICCGIGII ICGIAAAGIC IGGAAACGCG GAGICAGCG CCCIGCACCA IIAIGIICCG GAICIGCAIC GCAGGAIGCI GCIGGCIACC
                                                                                                                                                                                                                                                                                                                            II GII
                                                                                                                                                                                 cac8I
                                                                                                                          fnu4BI
                                                                                                                                        bsofI
                                                                                                                                                       bbvI
                                                                                                                                                                      sfani
                                                                                                                                                                                                                                                                                                                 bsrI
                                                                                                                                                                                   fokI
                                                                                                                                                                                                                                                                                                                             aciī
           mboi/ndeII[dam-]
                                                                                                                                                                                                                                                                                                                                            CIGGICCCGC CGCAICCAIA
                                                                                                                             mroI bsaBI[dam-]
                                                     dpnII[dam-]
                           mam [dam-]
                                        dpn [dam+]
                                                                    bstri/xholl
                                                                                    alwI[dam-]
                                                                                                                                                                                  accili[dam-]
                                                                                                                                                                                                                                                                                      sfani
Bau3AI
                                                                                                                                                         bspEI[dam-]
                                                                                                                                                                                                                                                                                                                 avall fnu4BI
                                                                                                                                                                                                                                                                                                   acil
                                                                                                                                                                                                                                                                                                                                 DSOFI
                                                                                                                                           bspMII
                                                                                                                hpall
                                                                                                                                                                         bsaWI
                                                                                                                                                                                                                                                            acli
                                                                                                                                                                                                                                                                         DSMFI
                                                                                                                                                                                                                                                                                      sau96I
                                                                                                                                                                                                                                                                                                   nlaIV
                                                                                                                                                                                                                                                                                                                                asuI
                                                                                                                                                                                                                                                                                                                                               3901 CIGIGGAACA CCIACAICIG IAITAACGAA GCGCIGGCAI IGACCCIGAG IGAITITICI
                                                                                                                                                                                           mslI
                                                                                                                                                                             hhaI/cfoI
                                                                                                                                                               fauDII/mvaI hiaPI
                                                                                                                                                                                            haeII
                                                                                                                                                                                                                                                                                                                                     ddeI
                                                                                                                                                                              bstul
                                                                                                                                      acil
                                                                                                                                                  thaI
                                                                                                                                                                                                                                                                                                           hhaI/cfoI
                                                                                                                                                                                                                                                                              cac8I
                                                                                                                                                                                                                                                                                                                                     eco47III
                                                                                                                                                                                                                                                                                           hinPI
                                                                                                                                                                                                                                                                                                                         haeII
                                                                                                                                                                                                                                                                                                                           tru9I
                                                                                                                                                                                                                                                                                                                                        nseI
                                                                                                                                                                         I loqu
                                                                                                                                                                                        bpuAI
                                                                                                                                                                                                                      3801
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4001 CAACGIICCA GIAACCGGGC AIGIICAICA ICAGIAACCC GIAICGIGAG CAICCICICI CGIIICAICG GIAICAIIAC CCCCAIGAAC AGAAAIICCC GIIGCAAGGI CAIIGGCCCG IACAAGIAGI AGICAIIGGG CAIAGCACIC GIAGGAGAGA GCAAAGIAGC CAIAGIAAIG GGGGIACIIG ICIIIAAGGG apol nlaIII mlI fokI sfaNI maeIII dsav nlaIII cauli hpall ncil Idem bslI bari maell

nspī

SCIFI

acil 4301 AAAACCTCTG ACACATGCAG CTCCCGGAGA CGGTCACAGC TTGTCTGTAA GCGGATGCCG GGAGCAGACA AGCCCGTCAG GGCGCGTCAG CGGGTGTTGG TITIGGAGAC IGIGIACGIC GAGGGCCICI GCCAGIGICG AACAGACAII CGCCIACGGC CCICGICIGI ICGGGCAGIC CCGCGCAGIC GCCCACAACC CICGACCIGO GCCIACIIGI COGICTGIAG ACACTIAGOG AAGIGCIGGI GCGACIACIO GAAAIGGOGI CGACGGAGOG CGCAAAGCCA CIACIGCCAC GCTGCCTCGC GCGTTCGGT GATGACGGTG muli maeili acii bsli nlaili acii melii meeili bpmi/gsul[dcm-] 4101 CCTTACACG AGCCATCAAG TGACCAAAAA GCCCTTAACA TGGCCCGCTT TATCAGAAGC CAGACATTAA CGCTTCTGGA GAAACTCAAC GGAAIGIGCC ICCGIAGIIC ACIGGIIIGI CCIIIIIIGG CGGGAAIIGI ACCGGGCGAA AIAGICIICG GICIGIAAII GCGAAGACCI CIIIGAGIIG hinPI nspBII fauDII/mvnI bstUI acil hphI bsh1236I hhaI/cfoI fauDII/mval hgaI fauDII/mvaI thaI mnli bsh1236I hhaI/cfoI **bsh1236I** bstul hinPI thaI bstul thaI tru9I fnu4HI **bsoFI** acil bbvI nspBII aluI IImd fnu4BI **bsoFI** bcgI bbvI GAGCIGGACG CGGAIGAACA GGCAGACAIC IGIGAAICGC IICACGACCA CGCIGAIGAG CIIIACCGCA drdI cauli scrFI hpall aluI ncil Idsm fokl dsav haeIII/palI sfani gau96I cac8I aciı asuI mslI tru9I mseI aluI asp700 maeIII hinfi tfii Idab esp3I bamBI bamAI hpall scrFI Idem Cauli dsav ncil nspar alui bali fpu4BI **bsoFI** bbvI nlaIII fnuDII/mvnI Idsu **bsh1236I** aluI hgaI fokI bstul acil thaI 4201

mplI

hgal drdI taqI

sfani

acii

cac8I

acil fnu4Bi bsoFI

bslI

bstNI

dsaV

apyl[dcm+] haeIII/pall

haeIII/palI

4701 AAGGCCAGGA ACCGTAAAAA

bstui bsh1236I

ecoRII

nlaIV

hgial/aspHI bsp1286 bsiHKAI bmyI ndeI apaLI/snoI alw44I/snoI AGAGTGCACC	ifoI mcrI baiEI cGGTCGTTCG GCCAGCAAGC	bsli cacsi haeIII/pali haei AGGCCAGCAA TCCGGTCGTT	
ddeI rsaI csp6I GCA GATTGTACTG A	hinpI hhal/cfoI fnu4HI pleI bsoFI mcr hinfI bbvI bsi GAC TCGCTGCGCT CGG	nlaili nspi nsphi afilii ggaaagaca igigagcaaa ccitictigi acactcgiti	
sfaNI fnu4HI bsoFI aciI TATGCG CATCAGA	mboli earl/ksp6321 hinpl sapi hinpl hinpl hal/cfol plei bsofi haeli acii mnli hinfi bbvi haelt cccrrccrc ccrcrcsc rccrcccrrccrrcrc rcccrccc	ATAACGCA GGAAAGP	
hgiAl/al bsp1286  sfaNI stant ddel bmyl bs bibkal  II fnu4HI ddel bmyl brokl acil acil acil bryl acil acil acil acil acil acil acil aci	mboli earl/ksp6321 hhal/cfol sapl hinPI hinPI sfaNI hhal/cfol pleI bsoFI mcrI acil haell acil mnli hinfi bbvi bsiEl artscccarc agcccrctr cccrtccrc ccrccrcac rccrccaca cccrccaca rarcccaca rccccaca cccccaca cccccaca rarccccaca rccccaca cccccaca cccccaca rarccccaca rccccacaca rarccccaca rccccacaca rarccccacaca rccccacacaca	nlaili bali cac81 haelil/pa gGCGGTAATA CGGTTATCCA CAGAATCAGG GGATAACGCA GGAAAGAACA TGTGAGCAAA AGGCCAGCAA CCGCCATTAT GCCAATAGGT GTCTTAGTCC CCTATTGCGT CCTTTCTTGT ACACTCGTTT TCCGGTCGTT	
bs acil ac scgar AgcgcagtG	sfaNI acii TAAGGAGAA ATACCGCATC	TAATA CGGTTATCC ATTAT GCCAATAGG	
			fauDII/mvaI
fou48I bsoFI bbvI hinPI nlaIII bsrI bs hhal/cfoI tth1111/as 4401 CGGGTGTCGG GCGCAGCCA TGACCCAGTC	acil sfani 4501 ATATGCGGTG TGAAATACCG CACAGATGCG TATACGCCAC ACTTTATGGC GTGTCTACGC	fnu4HI bsoFI acil fnu4HI acil bsoFI bsrBI bbvI cac8I 4601 GCTGCGGCGA GCGCTATCAG CGACGCCGCT CGCCATAGTC GAGTGAGTTT	FI thaI
4401 CGGGIGTC GCCCACAG	acil 4501 ATATGCGG	fnu4HI bsoFI acil fnu4HI bsoFI bsr bbvI cac8I bbvI cac8I cacGCGCGA	SCLFI mval

AAGGCCAGGA ACCGTAAAAA GGCCGCGIIG CIGGCGITIT ICCATAGGCI CCGCCCCCCI GACGAGCAIC ACAAAAAICG ACGCICAAGI CAGAGGIGGC IICCGGICCI IGGCAITITI CCGGCGCAAC GACCGCAAAA AGGIAICCGA GGCGGGGGA CIGCICGIAG IGITITIAGC IGCGAGIICA GICICCACCG

hhal/cfol hinPI

rmaI maeI bfaI

bsli haeIII/pall

haeI

scfI

acil

moli

	н н	ä
acii ACCIGICGGC IGGACAGGCG	hgial/aspHI bspl286 bsiHKAI bmyI apaLI/snoI alw44I/snoI TGTGCACGAA ACACGTGCTT	alwNI[dcm-] nu4HI soFI HI I maeIII bvI bsrI cA GCCACTGGTA
I hpall bawi cttacccat	aluI AGCTGGGCTG TCGACCCGAC	alwni fnu4HI bsoFI by I bsrI bbvI ACTGGCAGCA GC
scifi  scifi mval  mval  ecoRII dsaV  dsav bstNI  bstNI  apy1[dcm+] bssSI  tali mbl/cfoI  bsofi bsaMi  acic mspI  fnu4HI hpaII  acic mspI  bstNI  apy1[dcm+] bsaJI  alul mbl /cfoI  bsofi bsaMI  acic mspI  fnu4HI hpaII  acic dsaMI  acic mspI  fnu4HI hpaII  acic mspI  fnu4HI hpaII  acic dsaMI  acic dsaMI  acic mspI  fnu4HI hpaII  acic dsaMI  acic mspI  for daMI  for daMI  criticacic corrected accorded accor	hgiAI/aspl bsp1286 bsp1286 bsiEKAI basiEKAI bnyI col alul scfl ddeI c rcaiagctca cgcrgraggt arctcagtrc ggrgraggtc gtrcgcrcca agcrgggctg tgrgcacaa ig agali/sno lc alwi alw441/sn	fnu4HI bsoFI  mspI  pspBII  mcI bhori  mcri bbvi  bsiEI hhal/cfol hpaII  scccccctrc accccacc creccacc creccaccar creccaccar
bsli cfoi crccrgrrcc gaggaCaagg	GGTGTAGGTC	I I I GTAAGACACG
scrFI ecoRII hinPI apyl[dcm+] bssSI bsaJI alul mnll hhal/cfoI ccc rgGAAGCTCC CTCGTGCGCT CTC	ddeI ATCTCAGITC TAGAGICAAG	mspI hpall scrFl 'ncil pleI dsaV hinfI caulI GA GTCCAACCCG G
scrfi ecoRII apyl[dcm+] saJI alul m	scfl . CGCTGTAGGT	pl hi : ATCGTCTTGP
sc II dsav II dsav V bstNI ap II ap IGCM+) bsa	hinPI hhal/cfol haeli rgcccrttc rcatacctca accccaaag agtatcGagt	maelli mspl bsawi u hpall TCCGTAACT
scrFI mval ecoRII dsav bstNI apyI[d AGATACCAGG	hinPI hhal/cfoI haeII rGGGGCTTTC	fnu4HI bsoFI nspBII mspI acii hinPI mspI rI bbvI bsaWI tEI hhaI/cfoI hpaII ACCG CTGCGCCTTA TCCGGT
GAAACCCGAC AGGACTATAA CTTTGGGCTG TCCTGATATT	CTTTCTCCCT TCGGGAAGCG	fnu4B bsoFI nspBII aciI h mcrI bbvI bsiEI h ; AGCCCGACCG CTG
GAAACCCGAC	hinPI hhal/cf haell 4901 CTTCCCCT TCGGAAGCG TGCCGCTTI GAAAGAGGGA AGCCCTTCGC ACCGCGAAA	CCCCCCGTTC
4801	4901	5001

5101 ACAGGATTAG CAGAGCGAGG TATGTAGGCG GTGCTACAGA GTTCTTGAAG TGGTGGCCTA ACTACGGCTA CACTAGAAGG ACAGTATTTG GTATCTGCGC TGTCCTAATC GTCTCGCTCC ATACATCCGC CACGATGTCT CAAGAACTTC ACCACCGGAT TGATGCCGAT GTGATCTTCC TGTCATAAAC CATAGACGCG

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nlaIII
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                                                                                                                                                                                                                                                                                                     bapHI
                                                                                                                                                                                                                                                                                     rcal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AGITITIAAAT CAATCTAAAG TATATATGAG TAAACTTGGT CTGACAGITA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TCAAAATITA GITAGAITIC ATATATACIC AITIGAACCA GACIGICAAI
                                                                                                                                                                                                                                                                                                                    5301 ATTACGCGCA GAAAAAAGG ATCTCAAGAA GATCCTTTGA TCTTTTCTAC GGGGTCTGAC GCTCAGTGGA ACGAAAACTC ACGTTAAGGG ATTTTGGTCĀ
                                                                                                                                                                                                                                                                                                                                     TAAIGCGCGI CITITITICC TAGAGIICII CIAGGAACI AGAAAGAIG CCCCAGACIG CGAGICACCI IGCIITIGAG IGCAAIICCC TAAAACCAGI
                                                                                                                          AGACGACTIC GGICAAIGGA AGCCITITIC TCAACCAICG AGAACIAGGC CGITIGITIG GIGGCGACCA TCGCCACCAA AAAAACAAAC GIICGICGIC
                                                                                                           AGCGGTGGTT TITITGTTTG CAAGCAGCAG
                                           fnu4H]
                                                          DBOFI
                                                                            bbvI
                                                                                              cacel
                                                                                                                                                                                                                                                                      tru9I
                                                                                                                                                                                                                                                                                        mseI
                                                                                                                                                                                                                                                                                                         maelI
                                                                                                  acil
                                                                                                                5201 ICTGCTGAAG CCAGITACCI ICGGAAAAAG AGITGGTAGC ICTIGAICCG GCAAACAAAC CACCGCTGGT
                                                                                   nspBII
                                                                                                  acti
                                                                                                                                                                                                                                                                                                           hgal ddel
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ahaIII/draI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         tru9I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            mseI
                                                mbol/ndeII[dam-]
                                                                                    dpnII[dam-]
                                                                                                    alwi[dam-]
                                                                 dpnI[dam+]
                                                                                                                                                                                        mpol/ndeII[dam-]
                 hpaII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5401 TGAGATTAIC AAAAAGGAIC TICACCIAGA ICCITITAAA ITAAAAAIGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ACTCTAATAG TITITCCTAG AAGTGGATCT AGGAAAATIT AAITITTACT
Idsm
                                   sau3AI.
                                                                                                                                                                                                                                            dpnII(dam-
                                                                                                                                                                                                                          mpoll[dam-] dpul[dam+]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              tru9I
                                                                                                                                                                                                           mpol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ahaIII/draI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              nseI
                                                                                                                                                                          sau3AI
                                                                                                                                                                                                                                                                                                                                                                                                     mbol/ndell[dam-]
                                                                                                       aluI
                                                                                                                                                                                                                                                                              dpnII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           tru9I
                                                                                                                                                                                                                                                                                               alwI[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              mseI
                                                                                                                                                                                                                                                            dpnI[dam+]
                                                                                                                                                                                                                                                                                                                alwI[dam-] bstYI/xhoII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           dpnII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              bstYI/xhoII
                                                                                                                                                                                                                                                                                                                                                                                                                                          dpnI[dam+]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             alwI[dam-]
                                                                                                                                                                                            gau3AI
                                                                                                                                                                                                                                                                                                                                                                                          gau3AI
                                                                                                                                                                                                                                               mbol/ndeIl[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            mbol/ndeII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 alw1[dam-] bfaI
                                                                                                                                                                                                                                                                                                                                                                                                                             rmaI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               maeI
                                                                                                                                                                                                                                                                                 dpnII[dam-]
                                                                                                                                                                                                                                                                dppI[dam+]
                                                                                                                                                                                                                                                                                                    bstYl/xhoII
                                                                                                                                                                                                                                                                                                                                                                                                                                                             mpoll[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                dpnII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  bstYI/xhoII
                                                                                                                                                                                                                                                                                                                                                                                                                                            hphI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                dpn[dam+]
                                                                                                                                                                                                                              sau3AI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                sau3AI
                                                                                        maeIII
                                                                                                           ec 571 bsrI
                                                                                                                                                                                                                                                                                  fauDII/mval
                                                                                                                                                                                                                                                  hhaI/cfoI
                                                                                                                                                                                                                                   hinpi
                                                                                                                                                                                                                                                                                                        batuI
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5501 CCAATGCTTA ATCAGTGAGG CACCTATCTC AGGGATCTGT CTATTTCGTT CATCCATAGT TGCCTGACTC CCGTCGTGT AGATAACTAC GATACGGGAG

fokI

dpnII[dam-]

ddeI

dpn[[dam+]

ahdI/eaml105I

hinfi

pleI

mbol/ndeII[dam-]

hgici nlaIV

banI

tru9I

MDlI

sau3AI

**DSMAI** 

GGICCICCGA ICGIIGICAG AAGIAAGIIG GCCGCAGIGI IAICACICAI GGIIAIGGCA GCACIGCAIA fnu4BI **bsoft** bbvI nlaIII mslI haeIII/palI fpu4HI **bsoFI** acil eaeI cfrI mbol/ndell[dam-] mpli dpull[dam-] dpn1[dam+] pvul/bspCI gau3AI baiEI mcrI sau96I avall

TACAACACGT TITITCGCCA AICGAGGAAG CCAGGAGGCT AGCAACAGIC TICAITCAAC CGGCGICACA AIAGIGAGIA CCAAIACCGI CGIGACGIAI

5901 AIGIIGIGCA AAAAAGCGGI TAGCICCIIC

aluI

acil

mcri bsiEi hcgi fowl scal ddel acil bsrI scal acil bsrI scal acail brI scal acil bool attricttac torcotangar gcttictor gactogroad tactcaacca actcatroc acatacte according coloration acil
ddel TTCTG AGAA
ACCA AGTCA! IGGI ICAGI
rsal scal hI csp61 rgag TACTCA
rsal bsrI scaI maeIII hphI csp6I rgr GACTGGTGAG TAC?
ifani Sat Gctitic
foki i ca recgraad gr aggearre
f nlaiii TGTCATGCC ACAGTACGG
AITCICTIAC TAAGAGAAIG
6001

<u></u> 69/136	
sau3AI mbol/ndeII[dam-] dpnI[dam+] dpnII[dam-] bstXI/xhoII alwI[dam-] cGATC	
sau3AI mbol/nc dpnI[de dpnII[c bstXI/xi alwI[dau CTCAAGGATC GAGTTCCTAG	GCAAAACAG CGTITITGIC
hgal hinli/acyl hinli/acyl ahali/bsaHi hhal/cfol mspl hpall hpall hpall bspl286 cerFi bstUl bstUl ncil bshl2361 cauli hincli/hindli rfGCCGGCG rCAACACGCG AGAACTTTAA AAGTGCTCAT GTAACAGAAGCC CCGCTTTTGA GAGTTCCTAG	bgrI hgiAl/aspHI eco57I  sau3AI taqI bsp1286 mboIl[dam-]  mboI/ndeIl[dam-] bsiHKAI sau3AI sfaNI dpnI[dam+] apaLI/snoI mboI/ndeIl[dam-]  acil bstYI/xhoII maeIII bssSI dpnII[dam-]  acil bstXI/xhoII maeIII bssSI dpnII[dam-]  ATACCGCTGT TGAGATCCAG TTCGATGTAA CCCACTCGTG CACCAACTG ATCTTTACTT TCACCAGCGT TTCTGGGTGA AGACCCACT CGTTTTTGTC  AATGGCGACA ACTCTAGGTC AAGCTACATT GGGTGAGCAC TAGAAGTCGT AGAAAATGAA AGTGGTCGCA AAGACCCACT CGTTTTTGTC
maeli psp14061 xmni asp700 mboli GAAAA CGTICTICGG	hphI TCACCAGCGT AGTGGTCGCA
HI ma PSE XMDI ASP70C CATTGGAAAA GTAACCTTTT	-] am-] TCTTTTACTT AGAAATGAA
hgiAI/aspHI bsp1286 tru9I bsiHKAI mseI bmyI ahaIII/draI TTTAA AAGTGCTCAT CA	eco571 mboII[dam-] sau3AI sfaNI mboI/ndeII[dam-] dpnI[dam+] dpnI[dam-] tG ATCTTCAGCA TCT
tru9I mseI ahaIII/< AGAACTITAA A	hgial/aspHI bspl286 bsiHKAI si bmyI saal/snoI m alw441/snoI d iSI d
hinp! hhal/cfol thal fuuDil/mvnl bstUl bsh12361 cil CGC GCCACATAGC	hgia bspl bsiH bmyI apal alw4 I bssSI cccacrcGrG GGGTGAGCAC
hinpI hhal/ thal thuDII bstUI bshl23 aciI ATAATACCGC GC	bari sau3Ai taqi mboi/ndeli[dam-] dpui[dam+] alwi[dam-] stYi/xhoii maelii GATCCAG TTCGATGTAA CC
hgai hinli/acyi ahali/bsaHi pi ali ri ri ii li hincli/hindli cccc actacccc	bari sau3Ai ta mboi/ndeli[ dpni[dam+] dpni[dam-] alwi[dam-] bstYi/xholi TGAGATCCAG TTC
hinli/acyl hinli/acyl ahali/bsaHl hhal/cfol hgial/aspHI mspl hal/cfol hgial/aspHI maell dpn! hpall hal/cfol hgial/aspHI maell dpn! scrfi bstUl msel bmyl xmnl acil caull hincll/hindil acil 6101 TTGCCGGGG TCAACACGGG ATAATACCGC GGGTGTATCG TCTTGAAATT TTCACGAGTA GTAACCTTTT GCAAGAAACT CTCAAGGATC AACGGGCCGC AGTTTTGA GAGTTTTTAA AAGTGCTCATT GCAAGAAGCC CCGCTTTTGA GAGTTCCTAG	nspBII acii 6201 TTACCGCTGT AATGGCGACA
6101	CCT (BIN E 26)

eari/ksp6321 sspl 6301 GAAGGCAAAA TGCCGCAAAA AAGGGAATAA GGGCGACACG GAAATGTTGA ATACTCATAC TCTTCCTTTT TCAATATTAT TGAAGCATTT ATCAGGGTTA CTTCCGTTTT ACGGCGTTTT TTCCCTTATT CCCGCTGTGC CTTTACAACT TATGAGTATG AGAAGGAAAA AGTTATAATA ACTTCGTAAA TAGTCCCAAT acil fnu4HI bsoFI

TIOQU

SUBSTITUTE SHEET (RULE 26)

6401 TIGICICATG AGCGGATACA TAITIGAAIG TAITIAGAAA AATAAACAAA TAGGGGTTCC GCGCACATTI CCCCGAAAAG TGCCACCTGA CGTCTAAGAA AACAGAGTAC TCGCCTAIGI ATAAACTTAC ATAAATCTTI TTAITIGITI ATCCCCAAGG CGCGTGTAAA GGGGCTTTTC ACGGTGGACT GCAGAITCTI ahall/bsaHl aatii ddel hinlI/acyI maell nlaIV hhaI/cfoI fauDII/mvnI bsh1236I hinPI bstul thaI bspHI acil nlaIII rcal

sau96I
haeIII/palI
asuI mboII
ecol109I/draII
tru9I mnlI bpuAI
nseI

nlaIII

rcal

DSSSI DSPBI MSEI CATAGGCGTA TCACGAGGCC CTTCGTCTT CAA GEOT ACCATATATA AATAGGCGTA TCACGAGGCC CTTTCGTCTT CAA TGGATATTT TTATCCGCAT AGTGCTCCGG GAAGCAGAA GTT

FIG. 410

>length: 6563

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1119 1195 1425 1434 1446 1512 1695 1696 1752 2155 2375 2727 3002 3090 3339 3463
                                                                                                                                                                                                                                                                                                                                                                       2218 2233 2889 3292 4202 4259 4270 4319 4338 4619 4845 4935 4981 5238 5759 5859
                                                                                                                          3436 3448 3490 3544 3597 3613 3619 3700 3838 3967 3970 3981 4139 4155 4210 4266
                                                                                                  2628 2781 2784 2787 2906 2926 3005 3045 3094 3141 3226 3241 3309 3342 3367 3412
                                                                                                                                                 4442 4467 4505 4518 4544 4561 4604 4611 4632 4723 4751 4878 4897
                                                                                                                                                                                                                                                                                                                                                  72 121 252 320 398 532 589 648 1126 1144 1167 1325 1386 1906 2054 2075 2126
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                                                                           178 542 805 877 1340 1750 1826 2011 2039 2043 2182 2242 2384 2492 2501 2504
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            640 999 1347 1357 1449 1665 1713 1755 1764 2333 3262 3645 4705 4826 4839
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                                           1093 1963 4449
                                                                                                                                                                                                                                                                                                                                     ahdi/eam11051(GACNNNNGTC): 346 5566
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                                                                   1867 [dam-]
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645 6489
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                                                                                                                                                                                                                                                                                         ahall/bsaHI(GRCGYC):
                                                                                                                                                                                                                                                                                                                  ahalil/dral(TTTAAA):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           apall/snoI(GTGCAC):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   asp700(GAANNNTIC):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        apyI[dcm+](CCWGG):
                                                                                                                                                                                                                                                                                                                                                                                                                                                              alwI[dam-](GGATC):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           asp718(GGTACC):
                      acc651 (GGTACC):
                                                                      BCCIII (TCCGGA):
                                                                                                                                                                                                                                            aflii(ACRYGT):
 aatII(GACGIC):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   apol(RAATTY):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     apal (GGGCCC):
                                              accI(GTMKAC):
                                                                                                                                                                                                                                                                     ageI(ACCGGT):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   asuI (GGNCC):
                                                                                                                                                                                                                                                                                                                                                                  aluI(AGCT):
                                                                                               acii(CCGC):
                                                                                                                                                                                                                           acyl
```

Stop Template Primer

5' CAT GGT ATA GGT TAA ACT TAT TTA CAC 3' SL.97.2

NNS Randomization Primer

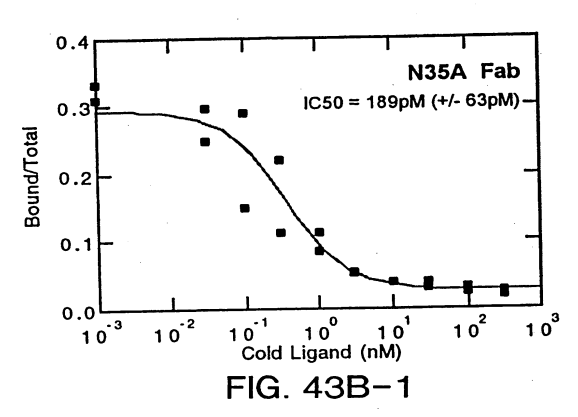
5' CAT GGT ATA GGT NNS ACT TAT TTA CAC 3' SL.97.3

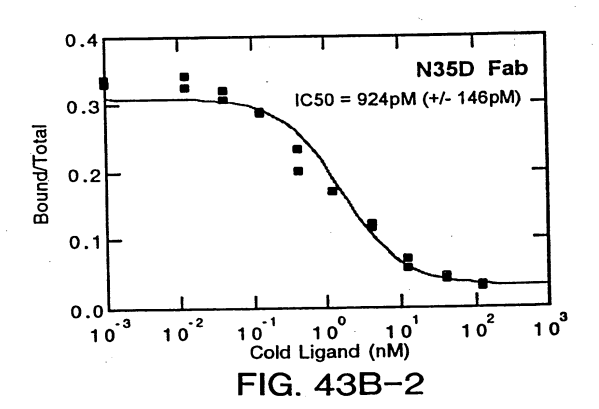
FIG. 42

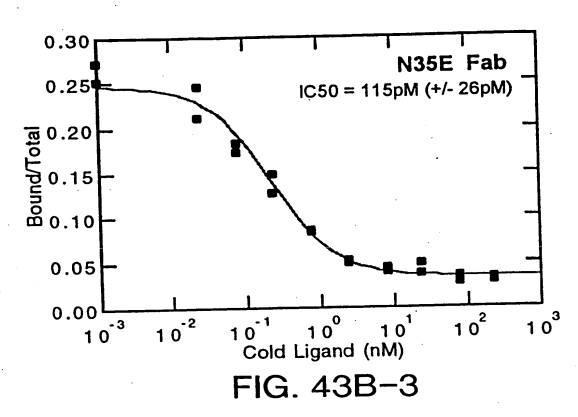
Randomization of Position N35 of Variable Light Chain CDR-1 Amino Acid Frequency

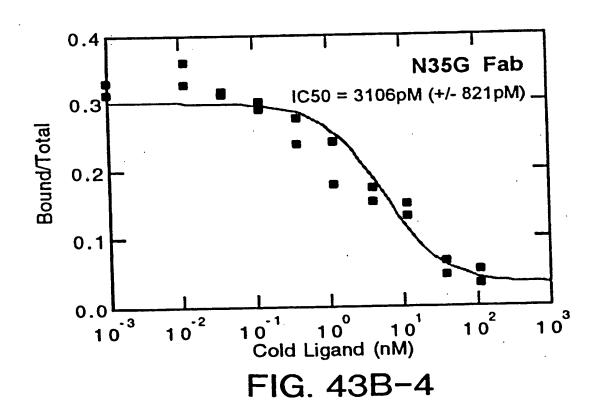
y) Sort #3	IC50 (nM)	4.9	3.1	3.1	0.1	0.2	NO	ND
don Libraı	% Total	5.6	16.6	16.6	22.2	5.6	5.6	1.9
y (NNS Co	Frequency % Total		9	33	4	2	₩	$\leftarrow$
Phage Display (NNS Codon Library) Sort #3	Amino Acid	Asparagine (wt)	Glycine	Aspartic Acid	Glutamic Acid	Alanine	Lysine	Serine

FIG. 43A

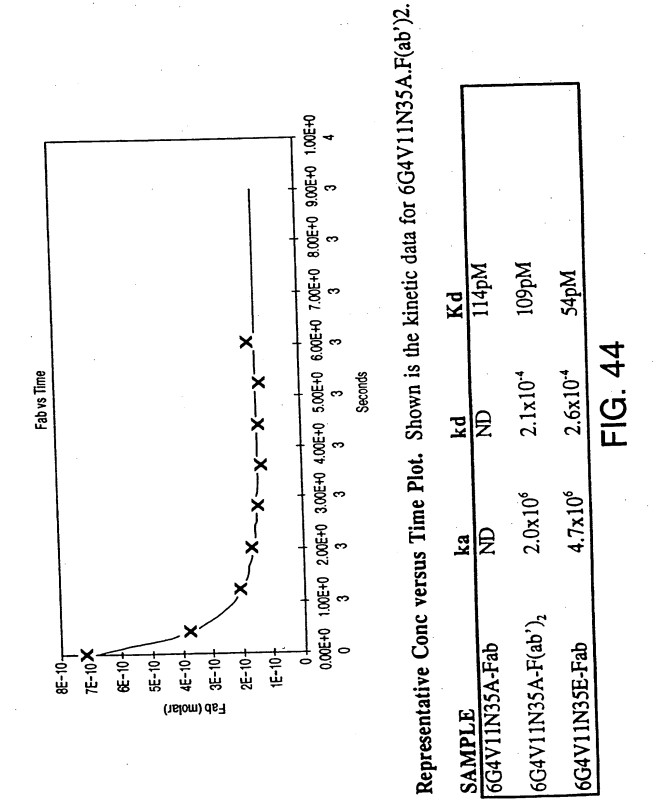








SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)

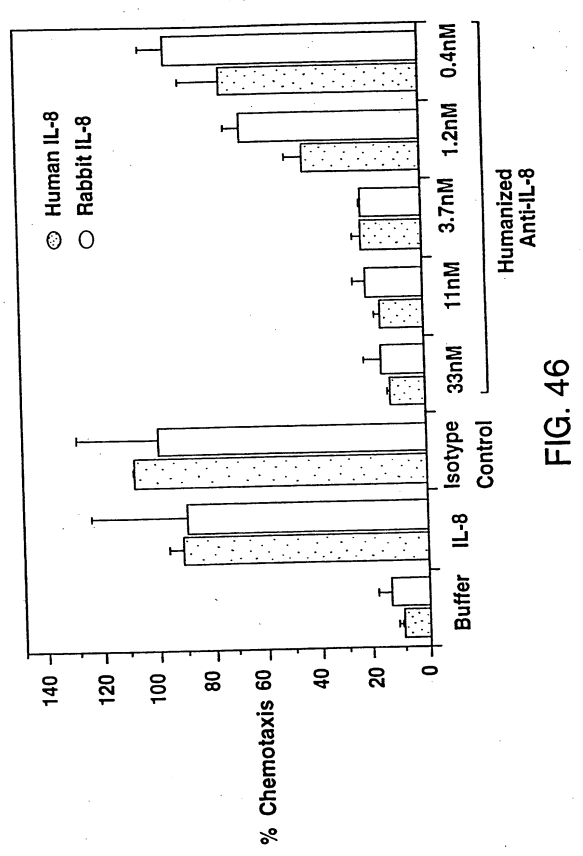
and the second second

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1 ATGAAAAAGA ATATCGCATT TCTTCTTGCA TCTATGTTCG TTTTTTCTAT TGCTACAAAC TACTTTTCT TATAGCGTAA AGAAGAACGT AGATACAAGC AAAAAAGATA ACGATGTTTG -23 M K K N I A F L L A S M F V F S I A T N 61 GCATACGCTG ATATCCAGAT GACCCAGTCC CCGAGCTCCC TGTCCGCCTC TGTGGGCGAT CGTATGCGAC TATAGGTCTA CTGGGTCAGG GGCTCGAGGG ACAGGCGGAG ACACCCGCTA -3 A Y A D I Q M T Q S P S S L S A S V G D 121 AGGGTCACCA TCACCTGCAG GTCAAGTCAA AGCTTAGTAC ATGGTATAGG TGAGACGTAT TCCCAGTGGT AGTGGACGTC CAGTTCAGTT TCGAATCATG TACCATATCC ACTCTGCATA 18 R V T I T C R S S O S L V H G I G E T Y 181 TTACACTGGT ATCAACAGAA ACCAGGAAAA GCTCCGAAAC TACTGATTTA CAAAGTATCC AATGTGACCA TAGTTGTCTT TGGTCCTTTT CGAGGCTTTG ATGACTAAAT GTTTCATAGG PGKAPKL LIY 38 <u>L. H.</u> W Y Q Q K 241 AATCGATTCT CTGGAGTCCC TTCTCGCTTC TCTGGATCCG GTTCTGGGAC GGATTTCACT TTAGCTAAGA GACCTCAGGG AAGAGCGAAG AGACCTAGGC CAAGACCCTG CCTAAAGTGA 58 N R F S G V P S R F S G S G T 301 CTGACCATCA GCAGTCTGCA GCCAGAAGAC TTCGCAACTT ATTACTGTTC ACAGAGTACT GACTGGTAGT CGTCAGACGT CGGTCTTCTG AAGCGTTGAA TAATGACAAG TGTCTCATGA 78 L T I S S L Q P E D F A T Y Y C S O S T 361 CATGTCCCGC TCACGTTTGG ACAGGGTACC AAGGTGGAGA TCAAACGAAC TGTGGCTGCA GTACAGGGG AGTGCAAACC TGTCCCATGG TTCCACCTCT AGTTTGCTTG ACACCGACGT 98 <u>H V P L T</u> F G Q G T K V E I K R T 421 CCATCTGTCT TCATCTTCCC GCCATCTGAT GAGCAGTTGA AATCTGGAAC TGCTTCTGTT GGTAGACAGA AGTAGAAGGG CGGTAGACTA CTCGTCAACT TTAGACCTTG ACGAAGACAA S G T 118 P S V F I F P P S D E Q L K 481 GTGTGCCTGC TGAATAACTT CTATCCCAGA GAGGCCAAAG TACAGTGGAA GGTGGATAAC CACACGGACG ACTTATTGAA GATAGGGTCT CTCCGGTTTC ATGTCACCTT CCACCTATTG 138 V C L L N N F Y P R E A K V Q W K 541 GCCCTCCAAT CGGGTAACTC CCAGGAGAGT GTCACAGAGC AGGACAGCAA GGACAGCACC CGGGAGGTTA GCCCATTGAG GGTCCTCTCA CAGTGTCTCG TCCTGTCGTT CCTGTCGTGG 158 A L Q S G N S Q E S V T E Q D S K D S T 601 TACAGCCTCA GCAGCACCCT GACGCTGAGC AAAGCAGACT ACGAGAAACA CAAAGTCTAC ATGTCGGAGT CGTCGTGGGA CTGCGACTCG TTTCGTCTGA TGCTCTTTGT GTTTCAGATG STL TLS KADY EKH KVY 178 Y S L S 661 GCCTGCGAAG TCACCCATCA GGGCCTGAGC TCGCCCGTCA CAAAGAGCTT CAACAGGGGA CGGACGCTTC AGTGGGTAGT CCCGGACTCG AGCGGGCAGT GTTTCTCGAA GTTGTCCCCT S P V T K S F N R G 198 A C E V T H Q G L S 721 GAGTGTTAAG CTGATCCTCT ACGCCGGACG CATCGTGGCC CTAGTACGCA ACTAGTCGTA CTCACAATTC GACTAGGAGA TGCGGCCTGC GTAGCACCGG GATCATGCGT TGATCAGCAT 218 E C Q

FIG. 45





SUBSTITUTE SHEET (RULE 26)

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5'-CTAGTGCAGTCTGGCGGTGGCCTGGTGCAGCCAGGGGGCTCACTCCGTTTGTCCTGTGCAGCTTCTGGCTACTCCTTC-3' N35AH1upr

N35AH1lwr

5'-TCGAGAAGGAGTAGCCAGAAGCTGCACAGGACAAACGGAGTGAGCCCCCTGGCTGCACCAGGCCACCGCCAGACTGCACT

AG-3'

Bold indicates nucleotide change destroying Pvull site.

CCGCCCCATG GCTGACTAAT TITITIATI

acil beaJI bell deal ncol

acil barl acil

acil

```
ANGENTECAT CICANITAGI CAGCANCCAG GIGIGGAANG ICCCCAGGGCI CCCCAGCAGG CAGAAGIAIG CAAAGCAIGC AICICAAITA
                                                                                                                                                                                                                                                                                                                                                                                                                                                    CITCAIACGI ITCGIACGIA GAGIIAAICA GICGIIGGIC CACACCIIIC AGGGICCGA GGGGICGICC GICIICAIAC GIITCGIACG IAGAGIIAAI
                                                                                                                                                                                                                                                                  1 TICGAGCICG CCCGACATIG AITAITGACI AGAGICGAIC GACACCIGIG GAAIGIGIGI CAGITAGGGI GIGGAAAGIC CCCAGGCICC CCAGGAGGA
AAGCICGAGC GGGCIGIAAC IAAIAACIGA ICICAGCIAG CIGICGACAC CIIACACACA GICAAICCCA CACCIITCAG GGGICCGAGG GGICGICGI
                                                                                                                                                                                                                                                       cac8I
                                                                                                                                                                                                                                                                                                                                                  nsii/avalii
                                                                                                                                                                                                                            apy1[dcm+]
                                                                                                                                                                                                                                                                                                                                                                                                            nspHI
                                                                                                                                                                                                                                                        nlaIV
                                                                                                                                                                                                                                                                                                                                                                                                                             cac8I
                                                                                                                                                                                                                                                                                                                                ppu10I
                                                                                                                                                                                ecoRII
                                                                                                                                                                                                              bstNI
                                                                                                                                                    SCLFI
                                                                                                                                                                                               dsav
                                                                                                                                                                                                                                          bsaJI
                                                                                                                                                                  mvaI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           nlallI
                                                                                                                                                                                                                                                           bsmFI
                                            >This has the pSvI backbone with the pRK7 cloning linker (pSvI7) and the intron DHFR(ID)
                                                                                                                                                                                                                                                                                                                                                                                                                               cacel
                                                            >mad from pSVI.WTSD.D by adding a linearization linker(LL) into the Hpal site
                                                                                                                                                                                                                                                                                                                                                                                                    apy1[dcm+]
                                                                                                                                                                                                                                                                                                                                                                                                                                 bsmFI nlaIV
                                                                                                                                                                                                                                                                                                                                                        ecoRII
                                                                                                                                                                                                                                                                                                                            SCLFI
                                                                                                                                                                                                                                                                                                                                                                                                                    bsaJI
                                                                                                                                                                                                                                                                                                                                                                                       batNI
                                                                                                                                                                                                                                                                                                                                                                        dsav
                                                                                                                                                                                                                                                                                                                                            mval
                                                                                                                                                           mbol/ndell[dam-]
                                                                                                                                                                                                                                                      nspBII
                                                                                                                                                                                                                                     pvull
                                                                                                                                             sau3AI aluI
                                                                                                                                                                                                                         hinfi taqi(dam-
                                                                                                                                                                                                                                                                                                                                                                                                                     apyI[dcm+]
                                                                                                                                                                                                       pleI dpnII[dam-]
                                                                                                                                                                          dpnI[dam+]
                                                                                                                                                                                          pvul/bspCI
                                                                                                                                                                                                                                                                                                                                                                           ecoRII
                                                                                                                                                                                                                                                                                                                                            SCLFI
                                                                                                                                                                                                                                                                                                                                                                                                         betNI
                                                                                                                                                                                                                                                                                                                                                                                         dsaV
                                                                                                                                                                                                                                                    beiEI
                                                                                                                                                                                                                                                                                                                                                             MVAI
        /home/ruby/vc/Immb1o/afan/ss.p6G425v11.N35A.choSD
                                                                                                                                                                                                                                         mcrI
                                                                                                                                                                                                                                          rmal
                                                                                                                                                                                                                                                        maeī
                                                                                                                                                                                                                                                                                                                                                                 nsil/avalli
                                                                                                                                                                                                                                                                                                                                    sfani
                                                                                                                                                                                                                                                                                                                                                 ppu10I
                                                                                                                                                                                                                                                                                                                                                                                 nlalli
> Wed May 7 18:27:36 1997
                                                                                                                                                                                                                                                                                                                                                                                                                              IHdeu
                                                                                                                                                                                                                                                                                                                                                                                                  sphi
                                           > length: 8120 (circular)
                                                                                                                                                                                  hgiAI/aspHI
                                                                                                                                                                                                                                                                                                                                                                                                                                                              101 GAAGTATGCA
                                                                                                                                                                                                 ecl136II
                                                                                                                                                                                                                 bsp1286
                                                                                                                                                                                                                                 DETHKAI
                                                                                                                                                                     hgiJII
                                                                                                                         aluI
                                                                                                                                                                                                                                                                banII
                                                                                                                                                                                                                                                   bmyI
                                                                                                                                          BBtI
                                                                                                                                                         Baci
```

CAGTCGTTGG INTCAGGGCG GGGATTGAGG CGGGTAGGGC GGGGATTGAG GCGGGTAAGA GCCGGGTAC CGACTGATTA AAAAAATAA

201 GTCAGCAACC ATAGTCCCGC CCCTAACTCC GCCCATCCCG CCCCTAACTC CGCCCAGTTC CGCCCATTCT

acil fokl

IXI	<b>3</b> / / / / 3 C	
/pali alli/e		
haeIII/palI uI mcrI eagI/xmaIII/eclXI eaeI cfrI bsiEI mspI hpaII GCTTATCCGG	14HI 5FI VI II nlaIII TG CCATCATGT AC GGTAGTACCA	rsal csp61 scal CAACTACTTC GTTCATGAAG
al rmar maer bfar nher cac81 alur CAAAAAGCTA	fnu4HI bsoFI bsoFI bbvI rsaI csp6I scfI mnlI aciI nlaIII GTACCGCCTA TAGAGCGATA AGAGGATTTT ATCCCGCTG CCATCATGGT CATGGCGGAT ATCTCCTAAAA TAGGGGCGAC GGTAGTACCA	xmnI asp700 GGAACGAGTT CCTTGCTCAA
rmal maeI styl bsaJI blaI avII[dam-] haeIII/palI stul mnll bfaI TTTGGAGGC TAGGCTTTTG	mnlI A AGAGGATTTT T TCTCCTAAAA	haeIII/pali haeI scrFI scrFI mval bsrBI ecoRII dsaV bstNI aclI bsal apyl{dcm+} bsal bsaJ mnli ddeI AGAACGGAGA CCTACCCTGG CCTCCGCTCA
rmal mael styl bsaJl blnI avrII stul haeI mnll bfaI TTTGGAGGCC TA	I TAGAGCGAT)	hae baerri mwar ecorii daav batur apyi[d i baaji i baaji r ccracccig
mnll beeri G AGGAGCTTT		bsmAI bsaI A AGAACGGAGA (
m b AGAAGTAGTG TCTTCATCAC	maell maelll AGTGACGTAA TCACTGCATT	. GGGATTGGC
el alul sali GAGCTATTCC	tfil hinfl ncil acil thai hpall flubil/mvnl dsaV caull CGGGAACGG TGCATTGGAA CGCGGATTCC CCGTGCCAAG GGCCCTTGCC ACGTAACCTT GCGCCTAAGG GGCACGGTTC	pflMI bsli sfani bemfi aactgcatcg tcgccgtgtc ccaaaatatg gggattggca ttgacgtagc agcggcacag ggttttatac ccctaaccgt
pali ddel ili ddel iaJi mnli al haeIII/pali CTCGGCCTCT GAG	tfii hinfi acii thai fnuDii/mvni bstii ccccGATTCC GCGCGATTCC	p1 bs bsmFI TCGCCGTGTC CO
fnu4HI bsoFI bsoFI bglI sfiI haeIII/palI mnlI mnlI mnlI bsaJI acil hGAGG CCGAGGCCGC CTCG	TGCATTGGAA	sfaNI AACTGCATCG TTGACGTAGC
fnu4HI bsoFI bsoFI bglI sfiI haeIII/palI mnlI mnlI ddeI mnlI mnlI haeIII/palI bsaJI mnlI mnlI bsaJI mcII haeIII/palI mnlI bsaJI aciI haeIII/palI stagagagagagagagagagagagagagagagagagagag	tfil boil ball ball ball ball ball ball ball cauli ccorrect Acceptage	pflMI bsli taqi sfaNi bsmFi 501 TCGACCATIG AGAGCGAGAGA AGCTGGTAAC TIGACGTAGC AGCGGCACAG GGTTTTATAC CCCTAACCGT TCTTGCCT
301	401	501

FIG. 48B

```
ahalll/dral
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         801 ACAACCGGAA TIGGCAAGTA AAGTAGACAT GGTTIGGATA GICGGAGGCA GIICIGITIA CCAGGAAGCC AIGAAICAAC CAGGCCACCI TAGACICITI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        IGITGGCCTT AACCGITCAI ITCAICIGIA CCAAACCIAI CAGCCICCGI CAAGACAAAI GGICCIICGG IACIIAGIIG GICCGGIGGA AICIGAGAAA
                                                        tru9I
                                                                                                                                                                                                                                                                                                                          701 AGGACAGAAT TAATATAGTI CICAGIAGAG AACICAAAGA ACCACCACGA GGAGCICATI TICITGCCAA AAGITIGGAI GAIGCCITAA GACITAITGA
                                                                                                                                                                                                                                                                                                                                           TGGTGGTGCT CCTCGAGTAA AAGAACGGTT TTCAAACCTA CTACGGAATT CTGAATAACT
                                                                       пвеI
                                                                                                   CAAAGAATGA CCACAACCIC TICAGIGGAA GGTAAACAGA AICTGGIGAI TAIGGGIAGG AAAACCIGGI ICICCAITCC IGAGAAGAAI CGACCIITAA
                                                                                                                  GITICITACT GGIGITGGAG AAGTCACCIT CCAITIGICI TAGACCACIA ATACCCATCC ITTIGGACCA AGAGGIAAGG ACICITCITA GCIGGAAAII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              hinfi
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ddel plel
                                                                                                                                                                                                                                                                                                aflii/bfri
                                                                                      ddel mboll tagl
                                                                                                                                                                                                                                                                                                                                                                                naeIII/pall
                                                                                                                                                                                                                                                                               tru9I
                                                                       hinfi
                                                                                                                                                                                                                                                                                                               fokI sfaNI mseI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               hinfI apyI[dcm+]
                                                                                                                                                                                                                                                                                                                                                                                                 haeI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                batNI
                                                                                                                                                                                                                                                                                                                                                                                                                                mvaI
                                                                                                                                                                                                                                                                                                                                                                                                                                                ecoRII
                                                                                                                                                                                                                                                                                                                                                                                                                 scrFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                dsav
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               nlaIII
                                                                            apy1[dcm+]
                                                                                                                                                                                                                                                                                                                   betXI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  tfil
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   apy1[dcm+]
                               ecoRII
scrFI
                                                              betNI
                mval
                                              daav
                                                                                            sexAI
                                                                                                                                                                                                                                                                                                                                                                                                                                                     ecoRII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 bstNI
                                                                                                                                                                                                                                                                                                                                                                                                                    BCLFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    dsaV
                                                                                                                                                                                                                                                                                                                                                                                                                                      nval
                                                                                                                                                                                                       hgiai/aspHI
                                                                                                                                                                                                                       ecl136II
                                                                                                                                                                                                                                      bap1286
                                                                                                                                                                                                                                                     DELHKAI
                                                                                                                                                                                         hgiJII
                                                                                                                                                                                                                                                                                      mnli aluI
                                                                                                                                                                                                                                                                                                      bassi banii
                                                                                                                                                                                                                                                                        bmyI
                                                                                                                                                            sstI
                                                                                                                                                                            BacI
                                                                                                                                                                                                                                                                                                                       bseRI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       MDlI
                                                                                 hphI
                                                                                                                                                                                                                                                                                                                         ball
                                                                                                 alwni[dcm-]
                                                                                  hinfi
                                                                                                                                                                                                                                                                                                                                                         TCCTGTCTTA ATTATATCAA GAGTCATCTC TTGAGTTTCT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        accI nlaIII
                                                                                    ear1/ksp632I
                                                   eco57I
                                                                      IIoqu
                                                                                                                                                                                                                                                                                                 tru9I
                                                                                                                                                                                                                                                                                                                mseI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               mspI
hpaII
                                                                                                                          601
```

FIG. 48C

```
ball ddeI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TACGTAAAAA TATTCTGGTA CCCTGAAAAC GACCGAAATC TAGGGGAACC GAAGCAATCT TGCGTCGATG TTAATTATGT ATTGGAATAC ATAGTATGTG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ACCCACCTAC AATTAATACA TAACCTTATG TATCATACAC
                                                                                                                                                                          GIGACAAGGA TCAIGCAGGA AITIGAAAGI GACACGIITI TCCCAGAAAI IGAITIGGGG AAAIAIAAAC CICICCCAGA AIACCCAGGC GICCICIG
                                                                                                                                                                                            CACTGTICCT AGTACGICCI TAAACTITCA CIGIGCAAAA AGGGICITIA ACTAAACCCC TITATATITG GAGAGGGICI TAIGGGICG CAGGAGAGAC
                                                                                                                                                                                                                                                                                                                                                                                          AGGICCAGGA GGAAAAAGGC AICAAGIAIA AGIIIGAAGI CIACGAGAAG AAAGACIAAC AGGAAGAIGC IIICAAGIIC ICIGCICCCC ICCIAAAGCI
                                                                                                                                                                                                                                                                                                                                                                                                            TCCAGGICCT CCTITITCCG TAGTICATAT TCAAACTICA GAIGCICITC TITCIGAIIG TCCTICIACG AAAGIICAAG AGACGAGGGG AGGATITCGA
                                                                                                                           ecoNI
                                 ahaII/bsaHI
                                                                     mn l I
                hinli/acyI
                                                                                                                                           apyI[dcm+]
                                                                                                                                                                                                                                                                                                                                                                           mnll
hgaI
                                                                                     ecoRII
                                                                                                                        bstNI
                                                                                                       dsav
                                                    BCLFI
                                                                     mval
                                                                                                                                                            bsaJI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                aseI/asnI/vspI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             tru9I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               mseI
                                                                                                                                                               mn]I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             fnu4HI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               bsoFI
                                                                                                                                                                                                                                                                                                                                                                            Ilodm
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                bbvI
                                                                                                                                                                                                                                                                                                                                                                                                                                 *END DHFR
                                                                                                                                                                                                                                                                                                                                                               sfaNI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ATGCATTITI ATAAGACCAT GGGACTITIG CTGGCTTTAG ATCCCCTTGG CTTCGTTAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          mbol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                                                                                mbol I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          bsaJI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               dpnII[dam-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             dpnI[dam+]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               alwI[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   bstYI/xhoII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            sau3AI
                                                                                                                                                                                                                                                                                                                                                                                 accI
                                                                                                                               maell
                                                                                                                                                afllII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    cacel
                                                                                                                                                                    maeIII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 dsal bsmFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               nlaIII
                                                                                                                                                                                                                                                                                                                                                                                    SfaNI
                                                                                                              mbol/ndeII[dam-]
                                                                                                                                                                    maelli alwi[dam-] apol
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   bsaJI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  styl
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ncol
                                                                                                                                                  dpnII[dam-]
                                                                           nlalII
                                                                                                                                  dpnI[dam+]
                                                                                                 sau3AI
                                                                                                                                                                                                                                                                                                                                 apyI[dcm+]
                                                                                                                                                                                                                                                                                                                                                                                     molI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     nsil/avallI
                                                                                                                                                                                                                                                                              ecoRII
                                                                                                                                                                                                                                                                                                              bstNI
                                                                                                                                                                                                                                           SCLFI
                                                                                                                                                                                                                                                                                               dsav
                                                                                                                                                                                                                                                            mval
                                                                                                                                                                                                                                                                                                                                                    sau96I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ppu101
                                                                                                                                                                                                                                                                                                                                                                     avall
                                                                                                                                                                                                                                                                                                                                                                                      asuI
                                                                                                                                                                                        901
                                                                                                                                                                                                                                                                                                                                                                                                       1001
```

7 48D

ecoRII

mval

SCIFI

sau96I avall asul

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mval fnu4HI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ecool091/drall
                                                                                                                                                            bstNI bsoFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 GOCAGGGGG TCACTCCGTT TGTCCTGTGC AGCTTCTGGC TACTCCTTCT CGAGTCACTA TATGCACTGG GTCCGTCAGG CCCCGGGTAA GGGCCTGGAA
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                                                     cauli
                          hpall
                                        dsav
             Idem
                                                                                                                                      Cauli
ncil
                                                                                               BCIFI
                                                                                                           ncil
                                                                                                                         dsav
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                                                                                                                                                                     cap61
                                                                                                                                                        rBal
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X
                                                                                                                                                                                                                                                                                   dsaI
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nlaIII alwI[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   GATCGATCGG GAATTAATTC GGCGCAGCAC CATGGCCTGA AATAACCTCT GAAAGAGAA CTTGGTTAGG TACCTTCTGA GGCGGAAAGA ACCATCTGTG
CTAGCTAGCC CTTAATTAAG CCGCGTCGTG GTACCGGACT TTATTGGAGA CTTTCTCCTT GAACCAATCC ATGGAAGACT CCGCCTTTCT TGGTAGACAC
                                                                                                                                                                                                                                    2801 AATAAAGCAA TAGCATCACA AATTTCACAA ATAAAGCATT TTTTTCACTG CATTCTAGTT GTGGTTTGTC CAAACTCATC AATGTATCTT ATCATGTCTG
                                                                                                                                                                                                                                                  TTATTICGIT ATCGIAGIGI ITAAAGIGIT TATTICGIAA AAAAGIGAC GIAAGAICAA CACCAAACAG GITIGAGIAG TIACATAGAA TAGIACAGAC
                                                                                                                             TGCAGCTTAT AATGGTTACA
                                                                                                                                          AGGGACAGAG GCCCATITAC TCACGCTGCC GGGATCTCAG CTGGACGTCT TCGAACCGGC GGTACCGGGT TGAACAAATA ACGTCGAATA TTACCAATGT
                                                                                   fnu4HI
                                                                                                    bsoFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           aclI
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                                                                                                                                 2701 ICCCIGICIC CGGIAAAIG AGIGCGACGG CCCIAGAGIC GACCIGCAGA AGCIIGGCCG CCAIGGCCCA ACTIGIIIAI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          asp718
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               haeIII/pall
                                                                                                                                                                                                                                                                                                                                                                                                                 nlaIV
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                                                                                                                                                                                                                                                                                                                                                                                                                                kpnI
Bau96I
                              asuI
                                             bsoFI nlaIII
                                                                                                         aluI haeIII/palI
                                                                                                                      hindili bgll beadi
                                                                            ncol
                                                                                           dsal
                                                              sfil styl
                               fnu4BI
                  actI
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                                                                            eaeI
                                                                                           cfrI
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                                                                                                                                                                                                                               bsml bfal
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                                                                               mael hincil/hindil
                                                                                                             begI
                                                                                              pstI
                                                                                                                           asul bfal acci bspMI
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                                                                rmal sall
                                                                                               sau96I hinfI
                                                   pleI
                                                                                                              haeIII/palI
                                                                                                                                                                                                                                                                                                                                                                                                                           hael
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                       hinPI
                                                                                                                                                                                                                                                                                                                    mbol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                              taq1[dam-] tru9I
                                                                                                                                                                                                                                                                                                                                                                                                                                             bspDI(dam-) mseI
                                                                                                                                                                                                                                                                                                                                                                                                                              clal/bsp106[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         dpnI[dam+] asp700
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          mpol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                            XmnI
                                                                                                                                                                                                                                                                                                                                                   dpnII[dam-]
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                                                                                       hpall
                                                                                                      dsav
                                            BCIFI
                                                          ncil
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                                                                                                                        bsmAI
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-IG. 48J

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BCFFI  BVBI  BCORII  BBENI  BEXAI  TTAGT CAGCAACCAG GTGTGGAAAG  AATCA GTCGTTGGTC CACACCTTC	acil foki ACTCC GCCCATCCCG CCCTAACTC FIGAGG CGGGTAGGC GGGGATTGAG	.I ddel   mnli alui mnli bseRI   sgccrcr GAGTATTCC AGAAGTAGTG
scrFI mval mval ecoRII ecoRII batNI apyI[dcm+] bsaJI bsmFI nlaIV cac8I cccAGGCTCC CCAGCAGGCA GAAGTATGCA AAGCATGCAT GAGTTATCA TCGTACTTCA AGCTTACTA GTCGTTGCT CTCAATTACT CACCAGCTCC CACACCTTC  TCAG GGGTCCGAGG GTCATTACT TCGTACGTA GAGTTAATCA GTCGTTGGTC CACACCTTTC	sfaNI  ppul01  nsil/avalli  lalli  hl  nspl  nspl  nspH  nspH  cac81  cac82  chargeatge atcteaatta greachae atagreege cectaacte griffed greatian cacated atagreege cectaacte	fnu4HI bsoFI bsoFI bglI stil ncol haelil/pall ddel muli bsaJI muli alui mi ssaJi scal ccccccc crccccrcr cacaratacacacccc crcccccrcr cacaratacacacacacacacacacacacacacacacaca
BOUTE BOLFI  BY BOUTE BOUTE  BY BOUTE BOUTE  BEAN BOUTE BOUTE  BOUT BOUTH  BOUT BOUT BOUTE  BOUT BOUTE  BOUT BOUTE  BOUT BOUTE  BOUT BOUT BOUTE  BOUT BOUTE  BOUT BOUT BOUT BOUTE  BOUT BOUTE  BOUT BOUTH  BOUT BOUTE  BOUT BOUT BOUTE  BOUT BOUTE  BOUT BOUTE  BOUT BOUT BOUTE  BOUT BOUT BOUTE  BOUT BOUTE  BOUT BOUTE  BOUT BOUTE  BOUT BOUTE  BOUT BOUT BOUTE  BOUT	nlalv scrfl mval nsil/avalli ecoRII dsav bstNI apyl(dcm+) bsaJl cac8l bsaJl 100 TCCCCAGGCT GCCTTCATAC GTTTCGTAGT TAGTCGTGG GGGTAGGC GGGGATTGAG AGGGTCCGA GGGTCGTCC GTCTTCATAC GTTTCGTAGT TAGTCGTG GGGTAGGC GGGGATTGAG AGGGTCCGA GGGTCGTCC GTCTTCATAC GTTTCGTAGT CACTCGTG TAGTCGTGG TAGTCGTTGG TAGTCGTGG TAGTCGTTGG TATCAGGCG GGGTTGAGG GGGGTAGGG GGGGTTGAG AGGGTCGTCC GTCTTCATAC GTTTCGTAGG TAGAGTTAAT CAGTCGTTGG TATCAGGCG GGGTTGAGG GGGGTAGGG GGGGATTGAG	fnu4HI bsoFI bsoFI styl ncol bslI dsal bslI dsal bsrI acil cccccatc cctcactar TITITIATT TATGCAGAGCCC CTCGCCCTCT GACTATTC GCGCCCAAG GCGGGGTAC CGACTGATA ANAMATAA ATACTCTC GGCTCGCCG GACCGGAGA CTCGATAAGG
3001 GAATGTGTGT CAGTTAGGGT GTGGAA	nlalv scrfl scrfl scrfl scorll dsav DstNl apyl(dcm+) bsayl AggggrccgA GGGTCGTCC G	3201

FIG. 48K

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GIACCGOCTA TAGAGICTAT AGGCCCACCC CCTIGGCTIC GITAGAACGC GGCTACAATT AATACATAAC CTITIGGATC GATCCTACTG ACACTGACAT
                                                                                                                                                                                                                                                                                                                                                                                                                      CATGGCGGAT AICTCAGAIA TCCGGGTGGG GGAACCGAAG CAATCTTGCG CCGATGTTAA TTATGTATTG GAAAACCTAG CTAGGATGAC TGTGACTGTA
                                                                                                                                                                  CGCGGATICC CCGIGCCAAG AGICAGGIAA
                                                                                                                                                                              TCCTCGBABA AAACCTCCGG ATCCGAAAAC GTTTTCGAT CGAATAGGCC GGCCCTTGCC ACGTAACCTT GCGCCTAAGG GGCACGGTTC TCAGTCCATT
                                                                                                                                                                                                                                           mbol/ndell[dam-]
                                                                                                                                                        hinfi
                                                                                                                                                                                                                                                                                                                                                                                                                                                      ~U2 match
                                                                                                                                                                                                                                                                                                               cla1/bsp106[dam-]
                                                                                                                                                                                                                                                                        dpnII(dam-)
                                                                                                                                                                                                                                                                                                                                                      mpol/ndell[dam-]
                                                                                                                                                                                                                                                           dpnI[dam+]
                                                                                                                                                                                                                                                                                     alwI[dam-]
                                                                                                                                                                                                                                                                                                                             bspDI[dam-]
                                                                                                                                                                                                                                                                                                    tadl[dam-]
                                                                                                                                                                                                                                                                                                                                                                                  dpnII [dam-]
                                                                                                                                                                                                                                 sau3AI
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                                                                                                                                                                                                                                                                                                                                                                                                                                        'removed ATG
                                                                                                                                                                                                                                                                                                                                                                                                 alwi[dam-]
                                                                                                                                                                                                                                                                                                                                            sau3AI
                                                                                                                               fnuDll/mvnI
                                                                                       hinfi
                                                                          tfil
                                                                                                                                                         bsh12361
                                                                                                       acil
                                                                                                                                             bstuI
                                                                                                                   thaI
                                                                                                                                                                                                                                                                                                                                                                                                   asel/asnl/vspl
                                                                                                                                                                        3301 AGGAGGCTIT ITIGGAGGCC TAGGCTTITG CAAAAAGCTA GCTTATCCGG CCGGGAACGG IGCATIGGAA
                                                                                                                                                                                                                                                                                                                                                                        fnuDII/mvnI tru9I
                                                                                          eagl/xmalII/eclXI
                                                                                                                                                                                                                                                                                                                                                                                        Bei
                                                                                                                                                                                                    ^seq from pSVI6B5-6G4VL: AvrII - HindIII frag
                                                                haeIII/palI
                                      hpall
                                                                                                                                                 mspl cauli
                                                                                                                                                                                                                                                                                                                                                                                                    bsh1236I
SCLFI
                        Idem
                                                     dsav
            ncil
                                                                                                                                                                                                                                                                                                                     fou4HI
                                                                                                                                                                                                                                                                                                                                   bsoFI
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                                                                                                                                   bsiEI
                                                                                                                                                                                                                                                                                                                                                             thaI
                                                                                                                                                             hpali
                                                                                                                                                                                                                                                                                                                                                                                                                                              ^sp6 promoter
                                                                                 BCLI
                                                                                                                      cfrI
                                                                                                          eael
                                                                                 aluI
                                                                                               rmal
                                                                                                                                                   cac8I
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                                                                                                                                      nheI
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                                                                                                            avrII[dam-]
                                                                                                                            haeIII/pall
                                            rmaI
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                                                                       styl
                                                                                                   blai
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lariat consensus^ IgG vH natural lariat restored^

FIG. 48L

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3701 ATAGGGTCAC CATCACCTGC AGGTCAAGTC AAAGCTTAGT ACATGGTATA GGTGCTACGT ATTTACACTG GTATCAACAG AAACCÁGGAA AÁGCTCCGAA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  CCACGAIGCA TAAAIGIGAC CATAGIIGIC IIIGGICCII IICGAGGCII
                                                                                                                                                                                                                                                                                                                ICINGINGCA ACTGCAACTG GAGIACAITC AGAIAICCAG AIGACCCAGI CCCCGAGCIC CCIGICCGCC ICIGIGGGCG
                                                                                       ball foki
                                                                                                                            GGTGAAAAAG AAAAAGAGGT GTCCACAGGT GAGGGTCCAG GTTGACGTGG AGCCAAGCGC TTCGATCGAA CCCGACGTAG CTAACTTAAG GTGGTACCCT
                                                                                                                CACCATGGGA
                                                                                                    beaJI
                                                                                                                                                                                                                                                                                                                                                                                                                                           apyI[dcm+]
                          nlallI
                                                                                                                                                                                                                                                                                          mnlI
                                                    pflMI
                                                                  ncol
                                                                                                                   GGGCTGCATC GALTGAATTC
                                                                                                                                                                                                                                                                                                      acil
                                                                                                                                                                                                                                                                                                                                                                                                     ecoRII
                                                                                                                                                                                                                                                                                                                                                                                                                              bstNI
                                                                              ecoRI
                                                                                                                                                                                                                                                                                                                                                                                                                  dsaV
                                                                                                                                                                                                                                                                                                                                                                                        mval
                                                                                           apol
                                                                                                       bbvI bspDI[dam-]
                                                                                                                                                                                                                         hgiAI/aspH
                                                                                                                                                                                                                                       ecl136II
                                                                                             bsoff tagi
                                                      clal/bsp106
                                                                                                                                                                                                                                                   bsp1286
                                                                                                                                                                                                                                                                 BETEKAI
                                                                                                                                                                                                                                                                                                                                ACCAGTACAT AGTAGGAAAA AGATCATCGT TGACGTTGAC CTCATGTAAG TCTATAGGTC TACTGGGTCA GGGGCTCGAG
                                                                                                                                                                                                               hglJII
                                                                                                                                                                                                                                                                                                        tth1111/aspI banII
                                                                                                                                                                                      sstI
                                                                                                                                                                                                  BacI
                                                                              fnu4HI
                                                                                                                                                                                                                                                                                              berI aval
                                                                                                                                                                                                                                                                                DemFI
                                                                                                                     3501 CCACTITITC ITITICICCA CAGGIGICCA CICCCAGGIC CAACIGCACC ICGGIICGCG AAGCIAGCII
                                                                                               bah1236I aluI
                                                                                 Cacel
                     rmal
                                                                                                           aluI
                                                                                                                                                  Aclouing linker
                                                                                                                                                                                                                                                                                                                                                                                                                      maell
                                                                                   betuI
                                  nael
                                                                                                                                                                                                                                                                                                                                                                                                                                               beaAI
                                                                                                                                                                                                                                                                                                                                                                                                                                    BnaBI
                                                           nheI
                                                                       fouDII/mvnI
                                                                                                             nruI
                                                                                                                                                                                                                                                                                                            ecoRV
                                                            thaI
                                                                                                             bsaJI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TGTACCATAT
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Bau96I
           avall
                          asuI
                                                                ecoRII
                                      BCTFI
                                                                                        betNI
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                                                    mvaI
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                                                                                                                    belI
                                                                                                                                                                                                                                                                                                       maeI
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sse8387I
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haeIII/palI
                                                                         fnu4HI
                                                                                  bsoFI
                                                                                             bbvI
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                                                                                                                    pstI
                                                                                                                                                                                                                                                              DBOFI
                                                                                                                                                                                                                                                                                                                                                                  GANATCIGGA ACTGCITCIG ITGIGIGCCI GCIGNAINAC ITCINICCCA GAGAGGCCAA
                                                                                                                                                                                                                                                                                 GATCAAACGA ACTGTGGCTG
                                                                                                                                                                                                                                                                                              CTAGITIGCI TGACACCGAC
                                                                                                                                                                                                                                                                        bbvI
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                                                                                                                                                                                                                                                                        dpnII(dam-)
                                                                                                                                                                                                                                                                                                                                                                              CITIAGACCI IGACGAAGAC AACACACGGA CGACITATIG AAGAIAGGGI
                                                                                                                                                                                                                                                             dpnI [dam+]
                                                                                                                                                                                                                                        sau3AI
                                                                                                                                                                                                                                                                                               GTOGGICITO IGAAGCGIIG AAIAAIGACA AGIGICICAI GAGIACAGGG CGAGIGCAAA CCIGICCCAI GGIICCACCI
                                                                                                                                                                                                                                                                                    3901 CAGCCAGAAG ACTTCGCAAC TTATTACTGT TCACAGAGTA CTCATGTCCC GCTCACGTTT GGACAGGGTA CCAAGGTGGA
                                                                                                                                                                                                                                                                                                                                                        asp700
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                                                                                                                                                                                                                                         hgici
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                                                     mbol/ndell[dam-]
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                                                                             dpnII[dam-]
                                                                                                                bstYI/xhoII
                                                                 dpnI[dam+]
                                                                                         alwi[dam-]
          hpall
                      belI
                                 DBBWI
Idsm
                                            sau3AI
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S
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D
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CCATGGCCCA ACTTGTTTAT TGCAGCTTAT AATGGTTACA AATAAAGCAA TAGCATCACA AATTTCACAA ATAAAGCATT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     GGIACCGGGT TGAACAAAIA ACGICGAAIA TIACCAAIGI TIATITICGII AICGIAGIGI IIAAAGIGII TAITICGIAA
                                                                                                                                                                                                                                                                                                                          4201 CTGACGCTGA GCAAAGCAGA CTACGAGAAA CACAAAGTCT ACGCCTGCGA AGTCACCCAT CAGGGCCTGA GCTCGCCCGT CACAAAGAGC TTCAACAGG
                                                                         4101 AGTACAGTGG AAGGTGGATA ACGCCTCCA ATCGGGTAAC TCCCAGGAGA GTGTCACAGA GCAGGACAGC AAGGACAGCA CCTACAGCCT CAGCAGCACC
                                   fnu4HI
                                                   ddel bsoFI
                                                              scfI mull bbvI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  sfani apol
                                                                                                                                                                                                                                                                                                                      maelil
                                                                                                                                                                              hgiai/aspHI
                                                                                                                                                                                                                                                   ddel cac8I
                                                                                                                                                                                          ec1136II
                                                                                                                                                                                                         bsp1286
                                                                                                                                                                                                                                                                                                        eco01091/drall
                                                                                                                                                                                                                        DBIHKAI
                                                                                                                                                                hglJII
                                                                                                                                                                                                                                                                   haeIII/pall
                                                                                                                                                                                                                                                                                           asul banil
                                                                                                                                                                                                                                                                              Bau96I aluI
                                                                                                                                                                                                                                         bmyI
                                                                                                                                       BBtI
                                                                                                                                                      Baci
                                                                                                                                                                                                                                                                                                                        alwi [dcm-]
                                                                                               TAGCCCATTG AGGGTCCTCT CACAGTGTCT CGTCCTGTCG
                                                                                                            0 0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       naelli
                                                               apy1 [dcm+]
                                                                                                                                                                                                                                                                                                                                                     GACTGCGACT CGTTTCGTCT GATGCTCTTT GTGTTTCAGA TGCGGACGCT
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                         ecoRII
                                                   bathI
                                      deav
BCLFI
             mvaI
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                                                                                                            TICCACCIAT IGCGGGAGGI
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dsal haelli/pall
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        CAGCAACCAG GIGIGGAAAG ICCCCAGGCI CCCCAGCAGG CAGAAGIAIG CAAAGCAIGC
                                                                                                                                                                                                                   TITITCACIG CATICIAGIT GIGGITIGIC CAAACICAIC AAIGIAICII AICAIGICIG GAICGAICGG GAATTAATIC GGGGGGGAC CAIGGGCTGA
AAAAAGIGAC GIAAGAICAA CACCAAACAG GITIGAGIAG IIACAIAGAA IAGIACAGAC CIAGCIAGCC CITAATTAAG CCGCGICGIG GIACCGGACT
                                                                                                                                                                                                                                                                                                                                                                                                     4501 AATAACCICT GAAAGAGGAA CITGGITAGG TACCTICTGA GGCGGAAAGA ACCAGCIGIG GAATGIGIGT CAGTTAGGGI GIGGAAAGIC CCCAGGCICC
                                                                                                                                                                                                                                                                                                                                                                                                                    TTATTGGAGA CITICICCIT GAACCAATCC ATGGAAGACT CCGCCITICI TGGTCGACAC CTTACACACA GTCAATCCCA CACCITICAG GGGTCCGAGG
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           mbol/ndell[dam-]
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                                                                                                                                                  mseI
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                                                                                                                   clal/bsp106[dam-]
                                                                                                                                                              mbol/ndell[dam-]
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                                           dpnII[dem-]
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                          dpnI[dam+]
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                                                          pvul/bspCI
                                                                                                      tagI[dam-]
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sau3AI
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                                                                                                                                                                                                                                                                                                                                                                    aaaaaataa atacgictcc gcotccggcg gagccggaga cicgataagg icticatcac icciccgaaa aaacciccgg atccgaaaac gititicgac
                                                                                    4701 ATCTCAATTA GTCAGCAACC ATAGTCCCGC CCCTAACTCC GCCCATCCCG CCCCTAACTC CGCCCAGTTCT CGCCCATTCT CGGCGGGTAC GGTGACTAAT TAGAGTTAAT CAGTCGTGG TATCAGGGCG GGGATTGAG CGGGATTGAG GCGGGATTAA TAGAGTTAAT CAGTCGTTGG TATCAGGGCG GGGATTGAG CGGGTAGGGC GGGGATTGAG GCGGGTTAAG GCGGGGTAAGA GGCGGGGTAC CGACTGATTA
nlaIII
                                                                         acil bsaJI
                                   ncol
                   styl
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                                                                  acil
                                                                                 acil fokl
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apyI[dcm+] ecoRII bathi dsav bsaJI 4901 TTACCICGAG CGGCCGCTIA ATTAAGGCGC GCCATTTAAA TCCTGCAGGT AACAGCTIGG CACTGGCCGT CGTTTTACAA CGTCGTGACT GGGAAAACCC AATGGAGCTC GCCGGCGAAT TAATTCCGCG CGGTAAATTT AGGACGTCCA TTGTCGAACC GTGACCGGCA GCAAAATGTT GCAGCACTGA CCCTTTTGGG maell maelil haeIII/palI eael cfrI berI aluI bagi maelii sse8387I bspMI scfI pstI ahalll/dral "linearization linker inserted into Hpal site tru9I msel tru91 bsh1236I msel msel bssHII swal hhal/cfol cacel hinPI ascI tru9I paci barBI bsoFI paer71 bs1EI cfrI xhoI fnu4HI aval bsoFI taqī

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fnuDII/mvnI
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                                                                                                                                                                                                  bstUI scfI
                                                                                                                                                                                                                                                                                                                                                                                                             5101 AGCCIGAATG GCGAATGGCG CCIGATGCGG TATTTTCTCC TTACGCATCT GTGCGGTATT TCACACCGCA TACGTCAAAG CAACCATAGT ACGCGCCTG
                                                                                                                                                                                                                                                             TCGGACTIAC CGCTIACCGC GGACTACGCC ATAAAAGAGG AATGCGTAGA CACGCCATAA AGTGTGGCGT ATGCAGTTTC GTTGGTATCA TGCGGGGAC
                                                                                                  accecaates stigaattas cegaacetce tetagegegg aagcegtcga cegcattate getteteegg gegtegetag egggaagget tetcaacgea
                                                                                      5001 IGGCGITACC CAACITAATC GCCTIGCAGC ACATCCCCCC TTCGCCAGCT GGCGTAATAG CGAAGAGGCC CGCACCGAIC GCCCTTCCCA ACAGTIGCGT
                                                                                                                                                                                                             bsh1236I
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         mbol/ndell[dam-]
                                             dpnII(dam-)
                     dpnI[dam+]
                                                         pvuI/bspCI
sau3AI
                                                                                  DB1EI
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                                  haeIII/palI
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                                                                                                                                                                                                                                                                                                                                                                   hinpi haeli
                                                           mnll acil
                                                                                 ear1/ksp6321
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                                        cacel
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                                                                    fnu4H]
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FIG. 48R

CCCAAAAAC GGGTTTTTG	pleI hinfI GACTCTTGTT CTGAGAACAA	tru9I alui msei a GCTGATTTAA	91 I AGCCAACTCC TCGGTTGAGG	VI KI Boll TC CGCTTACAGA
nlalv hgici taqi bani mnli GTGCTTTACG GCACCTCGAC CACGAAATGC CGTGGAGCTG GGGTTTTTG	tru9I mseI C TTTAATAGTG G AAATTATCAC	tru9I mseI T TAAAAATG A ATTTTTAC	hgial/aspHI bsp1286 bsiHKAI bsiHKAI bmyI ddeI fnu4HI apaLI/snoI rsaI bsoFI tru9I alw441/snoI csp6I sfaNI mseI GTGCACTCTC AGTACAATCT GCTCTGATGC CGCATAGTTA AGCCAACTCC	hinpi hhal/cfoi mspi thal thal thal funDii/mvni scrFi bstui nspBii bsh1236i drdi cauli acii
	maell plei drdi hinfi maell IGACGTIGGA GTCCACGTTC ACTGCAACCT CAGGTGCAAG	haelli/pall cg gccratrgg .gc cggaraacc	e fn bs sfani ct gctctgatgc	di anga acmagnene
mspl hpall hpall hpall nael cfrl01/bsrFl bmyl maell cac81 trtctcccca cgtrcgccg cttccccgt cacgccct cacgcccc Gaaggccc Gaaggccca Gttcgaatta	maell plei drdi hinf ctr rgacgrigga g	TITI GCCGAITI LAAA CGCCIAAA	hgial/aspHI bsp1286 bsiHKAI bmyI ddeI saI apaLI/snoI csp6I GTGCACTCTC AGTACAATCT CACGTGAGAG TCATGTTAGA	hinPI hhal/cfoi thal fnuDII/mvnI bstUI nspBII bsh12361 acii hgal
nlalv hgiJil bap1286 bmyl banil segect CCCTTA	GIT ITICGCC CAA AAAGCGG	TAT AAGGA1 ATA ITCCCI		hi thi frii frii brii acii bai
nlalv hgiJil bsp128 bmyI banli AA ATCGGGGCT	TG ATAGACG AC TATCTGO	CT TTTGATT GA AAACTAA	maell psp14061 tru91 msel TTAACGITTA CAATTTTATG	
aluI ST CAAGCICT SA GIICGAGA	haeIII/pali asuj GGGC CATCGCCC	aval ICTC GGGCTATI AGAG CCCGATAA		hinpi fnu4Hi baoFi iii hhai/cfoi bbvi
mspl hpall nael cfrl01/bsrFl cac8l GCCG CTTTCCCC	maell hae. dralli sau9( bsaal asul cc cctactcccc	bsli bsli a ca acctati	nvni trugi msei I taacaaaa Aa attgtttt	
mspl hpali nael cfrl01/bsrFl maelI cac8l 5301 TTTCTCGCCA CGTTCGCCG CTTTCCCGT CAGCTCTAA AAAGAGCGGT GCAAGGGGCA GTTCGAGTT	maeli haelii/pali draili sau961 hphi bsaal asui 5401 TTGATTTGG TGATGGTTCA CGTAGTGGC CATCGCCCTG ATAGACGGTT TTTCGCCCTT AACTAAACC ACTACCAAGT GCATCACCG GTAGCGGGAC TATCTGCCAA AAAGCGGGAA	bsli bsri bsri acaacactca acctaatct gggctattct titgatttat ggtttgacct tgttgtgagt tgggatagag cccgataaga aaactaaata	thal fnuDII/mvnI tru9I apol tru9I msel bstUI msel apol bsh1236I sspl 5601 CAAAATITA ACGCGAATT TAACAAATA	maelli maeli bsri nla bsaAi tthlili/aspi
1 TTTCTCGCC	h 1 TTGATTTGG AACTAAACC	bsrI CCAAACTGG GGTTTGACC	tr ms apol 1 CAAAATTT GITTTAAA	<b>.</b>
530.	540	550	560	,

5701 GCTATCGCTA CGTGACTGGG TCATGGCTGC GCCCCGACAC CCGCCTACAC CCGCTGÁCGC GCCTGACGG GCTTGTCTGC TCCCGGCATC CGCTTACAGA CGGATAGCGAT GCACTGACC AGGGCCGTAG GCGAATGCCT CGAATAGCGAT GCACTGACC AGGGCCGTAG GCGAATGCT CGAATAGCGAT GCACTGACC AGGGCCGTAG GCGAATGTCT

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sfani mboli(dam-) alw441/snoi
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GCACAGGGGG AAIAAGGGAA AAAACGCCGI AAAACGGAAG GACAAAAACG AGIGGGICII IGCGACCACI TICAIITICI ACGACTICIA GICAACCCAC
                                                                                                  haeIII/palI
                                                                                                                                                        eco01091/drall
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                                                                                                                                                                                                                                                                                                                                                                             5901 TACGCCTATT TITATAGGIT AATGICATGA TAATAATGGI TICITAGACG TCAGGIGGCA CITITCGGGG AAATGIGCGC GGAACCCCTA ITIGITIAIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              6001 TITCIAAATA CATICAAATA TGTATCCGCT CAIGAGACAA TAACCCIGAT AAAIGCTICA ATAATAITGA AAAAGGAAGA GTAIGAGTAI TCAACAITIC
                                                                                                                                                                                                                                                                                                                                                                                               ATGCGGATAA AAATATCCAA TTACAGTACT ATTATTACCA AAGAATCTGC AGTCCACCGT GAAAAGCCCC TTTACACGCG CCTTGGGGAT AAACAAATAA
                                                                                                                                                                       5801 CAAGCIGIGA CCGICICCGG GAGCIGCAIG IGICAGAGGI TITCACCGIC AICACCGAAA CGCGCGAGGC AGIAITCIIG AAGACGAAAG GGCCICGIGA
GIICGACACI GGCAGAGGCC CICGACGIAC ACAGICICCA AAAGIGGCAG IAGIGGCITI GCGCGCICCG ICAIAAGAAC IICIGCIIIC CCGGAGCACI
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                                                                                                 nepHI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   bsoFI
                                                                                Ideu
                                                                                                                                                                                                                                                                                                                                                                 rcal
                                                                                                                     fnu4HI
                                                                                                                                     DBOFI
                                                                                                                                                          bbvI
                                                                                                                                                                                                                                                                                                                                                                tru9I
                                                                                hpali
                             BCLFI
                                                                                                    dsav
                                                ncil
                                                              Igem
                                                                                                                                                          maeIII bamAI
                                                                                                                       esp3I
                                                                                                                                        bamBI
```

	100	/ 136
sephi trugi meei ahaiii/drai tT TTAAGTTCT AA AATTCAAGA	rsal csp61 bsrI scal hphI maeIII AGTA CTCACCAGTC	sau3Al mbol/ndell[dam-] dpnl[dam+] dpnl[dam+] pvul/bspCl mcrl bsiEl crGACAACGA
I 061 bsp1286 tru91 bs14KAI mseI bmyI ahaIII/draI TTTCCAATG ATGAGCACTT TTAAAGTTCT AAAAGGTTAC TACTGGAA AATTTCAAGA	real cep61 scal hi atgact tegttgagta ct	haeIII/pall eael cfrI fnu4HI bsoFI acil rGCGGC CAACTTACTT
maeli psp14061 xmnl asp700 mboli CC CCGAAGAACG TTTTC	I ddeI AT ACACTATTCT CAGA TA TGTGATAAGA GTCT'	nlaiii CC ATGAGTGATA ACAC GG TACTCACTAT TGTG
sau3Al mbol/ndeIl[dam-] dpnI[dam+] alwI[dam-] bstxI/xhoII A GATCTTGG	scil csel coll coll coll coll coll coll coll co	fnu4HI bsoFI bbvI mslI nlaIII rargcagrgc rgccaraacc arga
Bau3Al nspBII E mbol/ndeIl(dam-) m dpnI(dam+) c bstXl/xhoIl bsrI dpnII(dam-) acil bst a CTGGATCTCA ACAGGGGTAA c T GACCTAGAGT TGTCGCCATT c	scrFI ncii mspi mspi hpali dsav hinll/acyi hgal cauli ahall/bsaHi GTGATGA CGCCGGCGAA (	nlaiii scatgaca gtaagagaat
beri bari maelii taqi srggg TTACATCGAA CTG	acil ncil thai thai thai fuuDII/mvnI hpail bstUI hinli/acyl acil acil ddel scal hphi mae. bstUI hinpi ahai/cfoi ahail/bahi bcgi bsiEi bsoFi ddel scal hphi mae. cgatacaca cgcatatat cccatatat cccctt ctcttagc cagacata tctatataga acttacta cagacaca cagacacat cagacacata cccataacat cagacacata cagacacacata cagacacacata cagacacacata cagacacacata cagacacacata cagacacacacacacacacacacacacacacacacaca	haeIII/pall mb eae! dpi cfrI bsoFI bbvI mslI nlaili bst bst bst tgrchtigg tagaargcga tggcargarg carretta atacgrcac argastratic targaarga gactgriger trigaargaarga
bassi 6201 CACGAC GTGCT	6301 GCTAT CGATA	6401 ACAGA

6501 TCGGAGGACC GAAGGAGCTA ACCGCTTTTT TGCACAACAT GGGGGATCAT GTAACTCGCC TTGATCGTTG GGAACCGGAG CTGAATGAAG CCATACCAAA AGCCTCCTGG CTTCCTCGAT TGGCGAAAAA ACGTGTTGTA CCCCCTAGTA CATTGAGCGG AACTAGCAAC CCTTGGCCTC GACTTACTTC GGTATGGTTT

nlaili alwi[dam-]

acil

aluI

Bau96I avall Insp mnlI

mbol/ndeII[dam-] aluI

nlaIV

sau3AI

mbol/ndell[dam-] sau3AI maeIII

nlaIII

dpnI[dam+] dpnII[dam-]

Idam

hpall

dpnI[dam+]

dpnii[dam-] bsaWi

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mbol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       dpnII[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     dpnI[dam+]
                                                                               asel/asnl/vspl
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    sau3AI
                                                                                                                                                                                                   DBMAI
                                                                                                                                                                                                                    bsaI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ITTAAAAGGA TCTAGGIGAA GAICCITITI GALAATCICA IGACCAAAAI CCCITAACGI GAGIITICGI ICCACIGAGC GICAGACCCC GIAGAAAAAA
                                                                                                                                                                                                                                                                                                                                                                    CTCGCGGTAT CATTGCAGCA CTGGGGCCAG ATGGTAAGCC CTCCCGTATC GTAGTTATCT ACACGACGG GAGTCAGGCA ACTATGGATG AACGAAATAG
GAGCGCCATA GTAACGTCGT GACCCCGGTC TACCATTCGG GAGGGCATAG CATCAATAGA TGTGCTGCCC CTCAGTCCGT TGATACCTAC TTGCTTTATC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ACAGATCGCT GAGATAGGTG CCTCACTGAT TAAGCATTGG TAACTGTCAG ACCAAGTTTA CTCATATATA CTTTAGATTG ATTTAAAACT TCATTTTAA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAATITICCI AGAICCACII CIAGGAAAAA CIATIAGAGI ACIGGIIIIA GGGAATIGCA CICAAAAGCA AGGIGACICG CAGICIGGGG CAICIITICI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         IGICIAGGGA CICIAICCAC GGAGIGACIA ATICGIAACC AITGACAGIC IGGIICAAAI GAGIAIAIAI GAAAICIAAC TAAAITITGA AGIAAAAAIT
                                                                                                                                                                                                                                6701 GACTGGATGG AGGCGGATAA AGTTGCAGGA CCACTTCTGC GCTCGGCCCT TCCGGCTGGC TGGTTTATTG CTGATAAATC TGGAGCCGGT GAGCGTGGGT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             nseI
                                                                                                                                                                                                                                                CTGACCTACC TCCGCCTATT TCAACGTCCT GGTGAAGACG CGAGCCGGGA AGGCCGACCG ACCAAATAAC GACTATTTAG ACCTCGGCCA CTCGCACCA
                                                                                              ACAATTAATA
                                                                                                           GCTGCTCGCA CTGTGGTGCT ACGGTCGTCG TTACCGTTGT TGCAACGCGT TTGATAATTG ACCGCTTGAT GAATGAGATC GAAGGGCCGT TGTTAATTAT
                                              tru9I
                                                               mseI
                                                                                                                                                                                  cfr101/bsrFI
                                                                                                                                                                                                                  bpmI/gsuI[dcm-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ahaIII/draI
                                                                                                                                                                                                   nlaIV hphI
                                                                                                                                                                 hpaII
                                                                                              6601 CGACGAGCGT GACACCACGA TGCCAGCAGC AATGGCAACA ACGTTGCGCA AACTATTAAC TGGCGAACTA CTTACTCTAG CTTCCGGGCA
                                                                                                                                                  Idem
              hpall
                                                                                cauli
Idsm
                                SCIFI
                                                                dsaV
                                               ncil
                                                                                                                                                                                                                                                                                                                                                                                                                                                             tru9I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               mseI
                                 aluI
                                                                                bfal
                                               rmaï
                                                                 maei
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ddeI
                                                                                                                                                                                                                                                                                                                                                           ahdi/eam11051
                                                                                                                                                                                                                                                                                                                                           hinfI
                                                     berI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            maeII
                                                                    tru9I
                                                                                      mseI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            tru9I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           mseI
                                                                                                                                                                                                                          hpaII
                                                                                                                                                                                                          mspI
                                                                                                                                                                                        haeIII/palI
                                                    avili/fspl
                     hha1/cfoI
                                                                                                                                                                       196nes
       hinPI
                                                                                                                                                         bglI
                                      matI
                                                                                                                                                                                                              asuI
                                                                                     psp1406I
                                                                                                                                                                                                                              hhaI/cfoI
                                                                       maeII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      maeIII
                                                                                                                                                                                                              hinPI
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                                                                                                                                                                                                                                                                                                                                                                  Ilum
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                rcal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            mbol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                haeIII/palI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       tru9I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      mseI
                                                                         cac8I bsrDI
                                                                                                                                                                                              Bau96I
                                                                                                                                                                                                               avall
                                         fnu4HI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               dpnII[dam-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            dpnI[dam+]
                                                          bsoFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               alwI[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 betYI/xhoII
                                                                                            bbvI
                                                                                                                                                                                                                                                                                                                                  sau96I
                                                                                                                                                                                                                                                                                                                                                   nlaIV
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ahalii/drai bfai mboli[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          sau3AI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ban mali
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              mbol/ndell[dam-]
                                                                                             sfaNI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       hgici
                                                                                                                                                                                                                                                                                                                                                                                                                                                        nlaIV
                                                                                                                                                                                                                                                                                                                     fnu4HI
                                                                                                                                                                                                                                                                                                                                   bsoFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Bau3AI hphI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       mbol/ndell[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  dpnII[dam-]
                                                                                                                                                                                                                                                                                                                                                     bbvI
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  alwI[dam-]
                                                                                ms]I
                                                                                                                                                                                                                    acil
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 mael
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                rmaI
                                                                                           maeIII
                                                                                                                                                                                                                                                                                                                                        funDII/mvnI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         dpnI[dam+]
                                                                                                                                                                                                                                                                                                                                                                                                                                           ddeI
                                                                                                                                                                                                                                                                                                                                                                                                                                                           sau3AI
                                                                                                                                                                                                                     fokI
                                                                                                                                                                                                                                                                                                      acil
                                                                                                                                                                                                                                                                                                                                                        batui
                                                                                                                                                                                                                                                                                                                       thaI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      nseI
                                                                                                                                                                                                                                                                                                                                                                                           6801
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mbol/ndell[dam-]
                                                       dpnII[dam-]
                                       dpnI[dam+]
                                                                                                                                                                                                                            GCTACCAACT CITITICCGA AGGTAACTGG CTICAGCAGA GCGCAGATAC CAAATACTGT CCTICTAGTG TAGCCGTAGT TAGGCCACCA CTICAAGAAC
CGATGGTIGA GAAAAAĞGCT ICCATIGACC GAAGTCGICT CGCGTCTAIG GITTATGACA GGAAGATCAC ATCGGCATCA ATCCGGTGGT GAAGTICTIG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    7401 AGITACCGGA TAAGGCGCAG CGGTCGGGGT GAACGGGGGG TTCGTGCACA CAGCCCAGCT TGGAGCGAAC GACCTACACC GAACTGAGAT ACCTACAGCG
                                                                         alwi[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                   7301 TCTGTAGCAC CGCCTACATA CCTCGCTCTG CTAATCCTGT TACCAGTGGC TGCTGCCAGT GGCGATAAGT CGTGTCTTAC CGGGTTGGAC TCAAGACGAT
                                                                                                                                                                                                                                                                                                                                                                                                                     agacategtg geggatgtat ggagegagae gattaggaca atggteaeeg aegaeggtea eegetattea geaeagaatg geeeaaeetg agttetgeta
                                                                                                                          ACCAGCGGTG GITTGTITGC CGGATCAAGA
                                                                                                                                         agtitectag aagaacteta ggaaaaaag acgcgcatta gacgacgaac gtitgtitit tiggtggcga iggteseeac caaacaaacg gectagtiet
         sau3AI
                                                                                                           hpali
                                                                                             Idem
                                                                                                                                                                                              haeIII/pall
                                                                                                                                                                                                                                                                                                                                                                                      hinfi
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ddeI
                                                                                                                                                                                                                 haeI
                                                                                                                                                                                                                                                                                                                                                     hpall
                                                                                                                                                                                                                                                                                                                                                                                       Cauli
                                                                                                                                                                                                                                                                                                     SCIFI
                                                                                                                                                                                                                                                                                                                                                                      dsav
                                                                                                                                                                                                                                                                                                                                   Idsm
                                                                                                                                                                                                                                                                                                                     ncil
                                                                                                             nspBII
                                                                                                                                                                                                                  bslI
                                                                                                                              CIGCIGCIIG CAAACAAAA AACCACGCI
                                                                                                               acil
                                                                                                                                                                                  rmal
                                                                                                                                                                                                 maeI
                                                                                                                                                                                                                   bfal
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                                                                                                                                                                                                                                                                                                                                                                                            bsrI
                                                                                                                                                                                                                                                                                     fnu4HI
                                                                                                                                                                                                                                                                                                     DBOFI
                                                                                                                                                                                                                                                                                                                       bbvI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           alw44I/snoI
                                                                                                                                                                                                                                                                                                                                         fnu4HI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     hg1AI/aspHI
                                                                                                                                                                                                                                                                                                                                                          alwNI[dcm-]
                                                                                                                                                                                                                                                                                                                                                                           bsrI bsoFI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        apaLI/snol
                                                                                                                                                                                                                                                                                                                                                                                            bbvI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    bsp1286
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      bsinkaI
                                                                  cacel
                                                                                                                                                                                                                      hhal/cfol
                                                                               fnu4HI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          bmyI
                                                                                                  DEOFI
                                                                                                                 bbvI
                                                                                                                                                                                                     hinPI
                                                                                                                                                                                                                                                                                                                                                                                             maelll
                                                 fnuDII/mvnI
                                                                                                                                                                                                                                     GCTACCAACT CTTTTCCGA AGGTAACTGG CTTCAGCAGA
                                                                                                                                   TGCGCGTAAT
                                                                                 bsh1236I
                                                                                                                  hhaI/cfoI
                                                                  batul
                                                                                                  hinPI
                                                                                                                                                                                                                       eco57I
                                mbol/ndell[dam-] thal
                                                                                                                                    7101 TCAAAGGATC TTCTTGAGAT CCTTTTTTC
                                                                                                                                                                                                        bsrI
                                                                                                                                                                                                                         maeIII
                                                                               dpnII[dam-] dpnII[dam-
                                                                  dpn1[dam+] dpn1(dam+)
                                                                                                                       betYI/xhoII
                                                                                                     alwI[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             hinpi beiEI
                                                                                                                                                                                                                                                                                                                                                                                              mnlI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            bbvI mcrI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        nspBII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           acil
sau3AI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              hhaI/cfoI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Enu4HI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           bsoFI
                                                  mbol/ndell[dam-]
                   mpoll[dam-]
                                                                                                     batYI/xhoII
                                                                                                                       alwi[dam-]
                                                                                                                                                                                                                                                                                                                                                                                                  acil
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            hpall
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               beaWI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Idem
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               maeIII
                                                                                                                                                                                                                                              7201
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FIG. 48W

TCAATGGCCT ATTCCGCGTC GCCAGCCCGA CTTGCCCCCC AAGCACGTGT GTCGGGTCGA ACCTCGCTTG CTGGATGTGG CTTGACTCTA TGGATGTCGC

	1037	130	
scrFI mvaI ecoRII dsaV bstNI bstNI I bsaJI G GGAGCTTCCA C CCTCGAAGGT	nlaIV 11 26 AGCTATGGA 3CC TCGGATACCT	TG TGGATAACCG AC ACCTATTGGC	1 /efol 121 ic aatacgcaaa 3GG ttatgcgttt
scri mva. ecol dsa binpi mnli hhal/cfoi alui apy caratcas accarcas gascrica	sfaNI GATGCTCGTC AGGGGGG CTACGAGCAG TCCCCCC	tfii hinfi TGCGTTATCC CCTGATTCTG TGGATAACCG ACGCAATAGG GGACTAAGAC ACCTATTGGC	sapi hinPi mboli hhal/cfol eari/ksp6321 mnli acii haeli gaggaaggg aagagggggaaa crcctrcgc trcrcgcgg tratgcgttt
mspl hpall fnu4KI bsll bsoFI acil bswW acil gcggacagg alcograag cgcagggag gcggacagg alcograag cgrccag	qi CGATTTTGT GCTAAAAACA	haeIII/pall scrFI mval bsli ecoRII dsaV bstNI haeII/pall nspli apyl[dcm+] haeI affIII v haeI cac8I v haeI cac8I affIII tcctgcctt trgctcaca fgttcttcc tgccttarcc cctgattct ggataaccg	fnu4HI bsoFI bbvI pleI hinPI hinfI hhal/cfoI rcGCGCGGCAGCGCAGCCCGCCAGCCGCAGCGCAGC
	ta mnli drdi hgai grcgggttc GCCACCTCTG ACTTGAGCGT CAGCCCAAAG CGGTGGAGAC TGAACTCGCA	:111/ s11 cm+]	fnu4HI bsoFI bbvI til mcrI tili bs1EI cccacc ccaccaccc
hinPI hhal/cfol haeli TGAGCATTGA GAAAGGGCA GGGAGAAAG	I II [dcm+] GTATCT TTATAGTCCT CATAGA AATATCAGGA	haelli/pali haelli/ fuudhi scrFi bsoFi acii ecoRii thai bsli dsav fuuDII/mvni bstNi bstUi nlaiv hael caacGcGcc TTTTTACGT TCCTGGCCTT GTTGCCCGG AAAATGCCA AGGACCGGAA	fnu4HI bsoFI bbvI cac8I ac1I aluI ac1I bsoFI 7801 TATTACCGCC TTTGAGTGAG CTGATACCGC TCGCCGCCAC ATAATGGCGG AAACTCACTC GACTATGGCG AGCGCGTCG
hinPI hhal/cfol haell 7501 TGAGCATTGA GAAAGGGCT ACTCGTAACT CTTTCGCGGT GCGAAGGGCT	scrFI mvaI ecoRII dsav bstNI apyI[d 7601 GGGGGAAACG CCTGGT	cac81 7701 AAACGCCAG CA	acii 7801 TATTACCGCC TT ATAATGGCGG AA

maeIII ACCICACICA TIAGGCACCC CAGGCITIAC ACTITAIGCT TCCGGCTCGT AIGTIGIGIG GAATIGIGAG CGGATAACAA TITCACACAG GAAACAGCIA IGGAGIGAGI AAICCGIGGG GICCGAAAIG IGAAATACGA AGGCCGAGCA TACAACACAC CITAACACIC GCCTATIGIT AAAGIGIGIC CITIGICGAI CCGCGCGTTG GCCGATTCAT TAATCCAGCT GGCACGACAG GTTTCCCGAC TGGAAAGCGG GCAGTGAGCG CAACGCAATT AATGTGAGTT GGCGGAGAGG GGCGCGCAAC CGGCTAAGTA ATTAGGTCGA CCGTGCTGTC CAAAGGGCTG ACCTTTCGCC CGTCACTCGC GTTGCGTTAA TTACACTCAA asel/asnl/vspl tru9I mseI hhal/cfol acil bsrBI cacel acil berI hpaII Idsm eael tfil asel/asnl/vspl cfrl hinfi msel nspBII aluI IInad tru91 haeIII/pall hgici apyi[dcm+] ecoRII SCIFI nlaIV bstNI dsav fnuDII/mvnI HVAI fnuDII/mvnI **bsh1236I** hhal/cfol Dsh1236I bstul hinPI bstul thaI bslI CCGCCTCTCC mnll acii 8001 7901

thaI

FIG. 48Y

asel/asnl/vspl

asp700

nlalII

XmnI

8101 TGACCATGAT TACGAATTAA ACTGGTACTA ATGCTTAATT

tru9I

msel

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3562 3566 3676 3733 3792 4270 4288 4311 4344 4554 4842 4896 4954 5047 5333 5590 5803 5822 6516 6579 6679 7200 7457 7593 7819 7937 8096
                                                                                                                                                             5166 5203 5217 5220 5248
                                                                                                                                                                                                               7166 7175 7310 7420 7541 7560 7687 7715 7806 7827 7834 7877 7901 7911 7967 8070
                                                                                                                                     3167 3179 3188 3200 3210 3221 3267 3372 3404 3449 3686 3949 4021 4318 4542 4727
                                                                                                                                                                                            6713 6804
                                                                                               823 1039 2738 4237
217 229 238 250 260 271 317 422 454 485 574 1385 1795 1871 2248 2250 2758 2982
                                                                                                                                                                                                                                                                                                                                                                                                                              5 44 332 386 390 753 1097 1165 1370 1431 1951 2603 2751 2784 3282 3336 3340
                                                                                                                                                                                         5751 5790 5979 6026 6125 6234 6311 6355 6476 6522
                                                                                                                                                                    4739 4748 4760 4770 4781 4827 4910 4914 5070 5127 5153
                                                                                                                                                                                                                                                                                                                                                            988 1690 1858 5117 5947 6329
                                                                                                                                                                                                                                                                                                                                                                                       696 4935 6290 6982 7001
                                                                                                                                                                                                     5275 5680 5699 5741
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1876 5651 6198 7444
                                                                     2969 3967 4529
                                                                                                                                                                                                                                                                                                                                                                                                                 ahdI/eam11051(GACNNNNGTC): 2087 6865
                                                                                                                                                                                                                                                            see hinli
                                                                                                                                                                                                                                                                                  786
932 7758
                                                                                                                                                                                                                                                                                                                                       1833
                                                                                                                                                                                                                                                                                                                                                                 ahall/bsaHI (GRCGYC):
                                                                                                                                                                                                                                                                                                                                                                                           shalli/dral(TTTAAA):
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         alw441/snoI(GTGCAC):
                                                                                                                                                                                                                                                                                    afili/bfri(cTTAAG):
                                                                                                                                                                                                                                                                                                                 aflil(ACRYGT):
                                                                              acc651 (GGTACC):
                                                 aatII(GACGTC):
                                                                                                                                                                                                                                                                                                                                           ageI (ACCGGT):
                                                                                                        accI (GTMKAC):
>length: 8120
                                                                                                                                                                                                                                                                                                                                                                                                                                                aluI (AGCT):
                                                                                                                                ac11(CCGC):
```

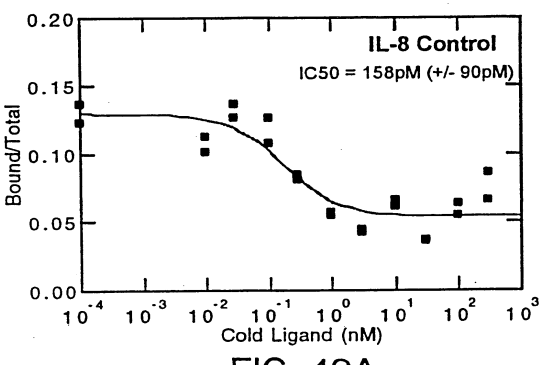


FIG. 49A

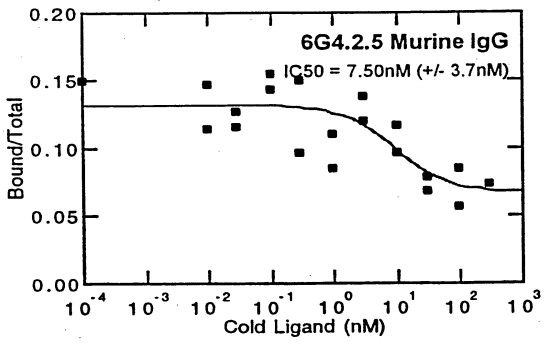


FIG. 49B

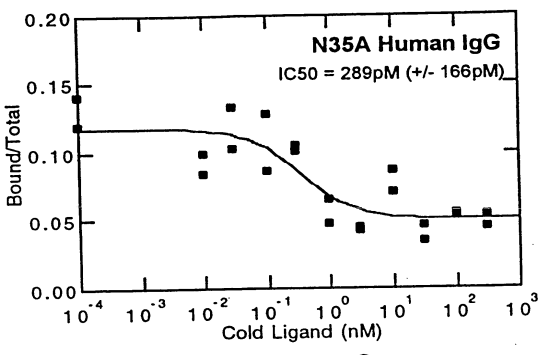


FIG. 49C

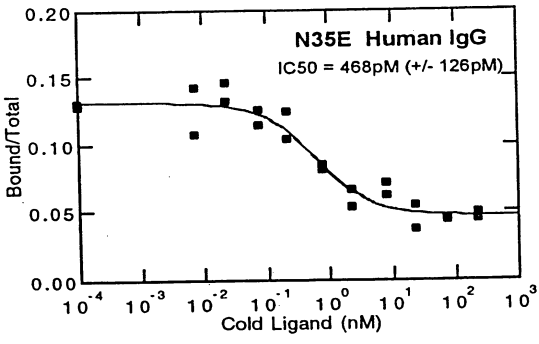
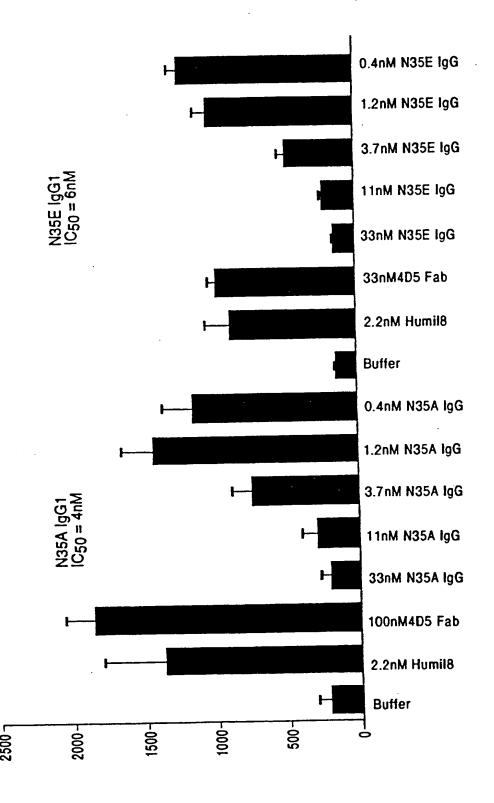
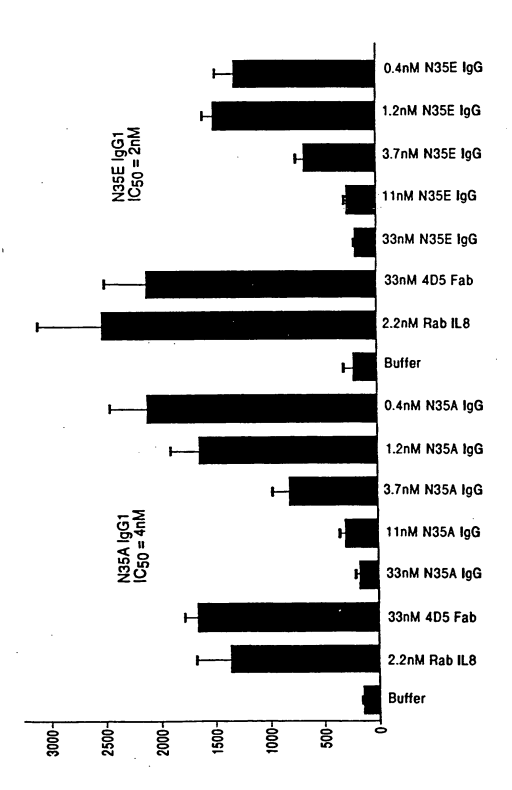


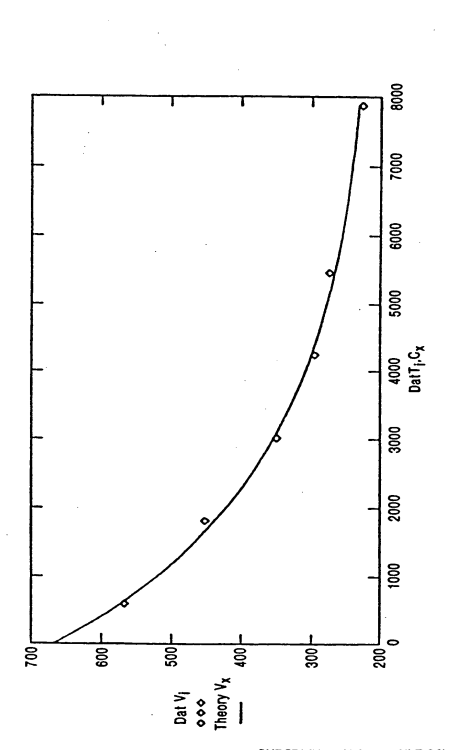
FIG. 49D



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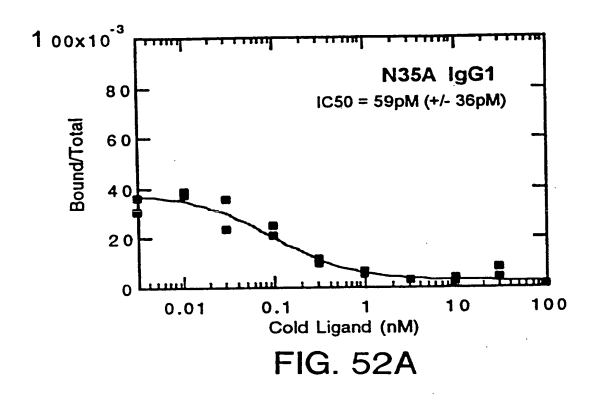


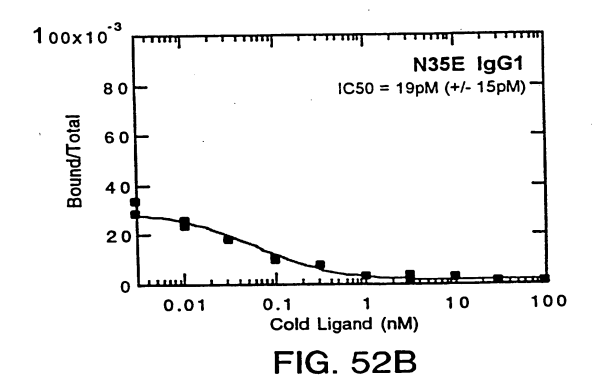




Representative Conc versus Time Plot. Shown is the kinetic data for 6G4V11N35A.IgG1

SAMPLE	ka	kd	Kd	
Murine 6G4.2.5 IgG2a	8.3×10 <sup>5</sup>	2.9×10-4	350pM	
6G4V11N35A-IgG1	$8.7x10^{5}$	$7.7x10^{-5}$	88pM	j
6G4V11N35E-IgG1	$3.0 \times 10^6$	1.4x10 <sup>-4</sup>	49pM	FIG. 5

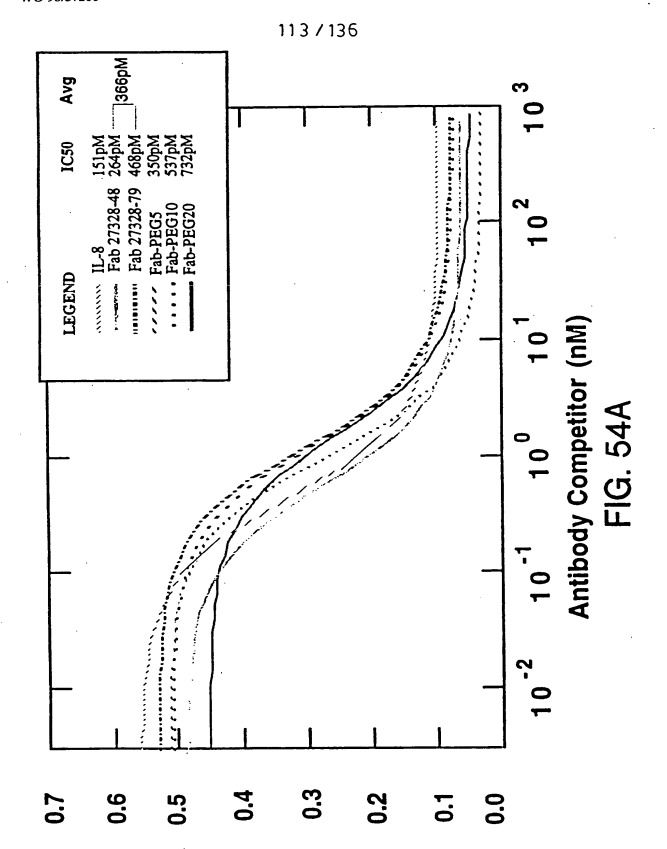




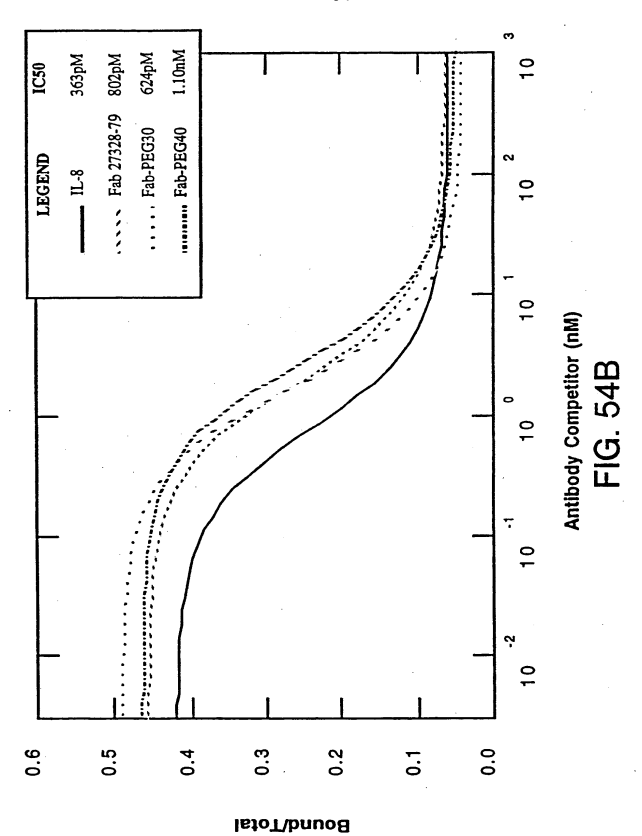
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781 -1	AA. TT	AAG PTC	GGT CCA	'AT TA	CTAG GATC	AGGT TCC1	MG AAC	AGG? TCC!	IGA? ACTI	AAAA	TAC	CTT	AAAG TTTC K	T	TATA TATA I	GCG'	TAA	AGA	<b>IGAA</b>	CGT
	AG.	ATA	CAA	GC	TTTT AAAA F	AAGI	ATA	ACG2	ATG:	TTTG	CG	CATY	CGA	/C	AGGT TCCA	AGT	CGA	AGTO TCAO V	CTC	:AGA
-11	S	M	F.	V	F.	S	1	A	1.	14	A	1	^		•	¥	_	•	¥	
901	GG	CGG	TGG	CC	TGGT	GCA	GCC	AGG	GGG(	CTCA	CT	CCG'	TTC	T	CCTG	TGC	AGC	TTCT	CGC	TAC
	CC	GCC	ACC	:GG	ACCA	CGT	CGG	TCC	CCC	GAGT	GAG	GGC	AAAC	:A	GGAC	ACG	TCG	AAG	YCCC	ATG
8	G	G	G	L	V	Q	P	G	G	S	L	R	L	S	С	A	A	S	<u>G</u>	Y
961	TC	CTT	CTC	GA	GTCA	CTA'	TAT	GCA	CTG	GGTC	CG	rca(	GCC	CC	CGGG	TAA	GGG	CCTY	GAZ	TGG
	AG	GAA	GAG	CT	CAGT	GAT	ATA	CGI	GAC	CCAG	GC	AGT	- CGG	فاد ت	GCCC	ATT	CCC	T	.C.I.I	W
28	S_	_F_	_S_	S_	Н	_Y	м	_н	W	V	K	Q	A	P	G	V	G	13	E	"
1021	GT	TGG	ATA	ΔTA	TTGA	TCC'	TTC	CAA'	TGG'	TGAA	AC'	TAC	GTA1	ra	ATCA	AAA	GTT	CAAC	GGG	CCGT
	CA	ACC	TAT	'nΤ	AACT	'AGG	AAG	GTT	ACC.	ACTT	TG	ATG	CATA	T	TAGT	TTT	CAA	GTT	CCC	GCA
48	v	G	Y	I	D	P	S	N_	G	E	T	T	Y	N	0	_K_	F	K	G	R
-	•	_	_																	
1081	Jel	CAC	Lalal	TAT	CTCG	CGA	CAA	CTC	CAA	AAAC	AC.	AGC.	ATAC	CC	TGCA	GAT	GAA	CAG	CTC	CGT
	AA	GTG	AAA	TA	GAGC	GCT	GTT	GAG	GTT	TTTG	TG	TCG'	TATO	<b>3</b> G	ACGT	CTA	CTT	GTC	GAC	CGCA
68		T					N			N	T	A	Y	L	Q	M	N	s	L	R
	_	_	_																	
1141	GC	TGA	.GG?	CA	CTGC	CGT	CTA	TTA	CTG	TGCA	AG.	AGG	GGAT	rr	ATCG	CTA	CAA	TGG'	IGAC	CTGG
	CG	ACI	CCI	T	GACG	GCA	GAT	AAT	GAC	ACGT	TC	TCC	CCT	AA	TAGO	GAT	GTT	ACC	ACTY	SACC
88	A	E	D	T	A	V	Y	Y	С	A	R	G	<u>D</u> _	<u>Y</u>	R	<u>Y</u>	_N	<u>G</u> _	D	<u>-M</u>
																		C3.C	~ 3 3/	
1201	TT	CTI	CG	\CG	TCTC	GGG	TCA	AGG	AAC	CCTG	GT	CAC	CGT	C'I'	CCTC	الفالفات	CIC	CAC		
•	AA	GAA	GCI	rgc	AGAC	CCC	AGT	TCC	TTG	GGAC	CA	GTG	GCA	GA.	GGAG	CCG	JAG.	T	21.16	C
108	F	F	D_	<u></u> Y	W	G	Q	G	Т	ь	V	T	V	5	S	A	3	•	K	G
					<b></b>		~~~	300	<b>C</b> MC		2.2	C D C	C N C (	сф	CTGG	ימכני	CAC	AGC	GGC	CTG
1261	CC	:A'I'C	:GG'	ICT	TCCC	CCT		ACC	CIC	CICC	MA	OAG OTO	CAC	- T	GACC		CTC	TCG	CCG	GAC
															GACC	G	.G.1.С.	A	A D	ī.
128	Þ	S	V	F.	P	ь	A	P	5	5	V	3	1	3	G	<b>G</b> .	•	**	••	_
1221	CC	יריתי		ייי	TC A	ACC N	בידים	Curi	ነገጋን	CGAA	CC	CGT	GAC	GG	TGTC	GTC	GAA	CTC	AGG	CGCC
1321	CC	C 10			yC.T.	ינטטני זייררייו	TAT	GAA	ന്ദ്ര	CCTT	GG	CCA	CTG	CC	ACAG	CAC	CTT	GAG	TCC	CCCC
149					K						P	v	T	v	S	W	N	s	G	A
140	. G	C	ם	•			•	•	•	_	•	•	_						• •	
1381	C	(GAC	CA	GCG	GCG'	rgca	CAC	CTI	CCC	GCT	GT	CCI	ACA	GT	CCTC	AGC	ACT	CTA	CTC	CCTC
	G	CTY	CTY	CGC	CGC	ACGT	GTG	GAA	GGG	CCGA	CA	GGA	TGT	CA	GGAC	STCC	TGA	GAT	GAG	GGAG
168	L	T	S	G	V	Н	$\mathbf{T}$	F	P	A	v	L	Q	S	s	G	L	Y	S	L
1441	A	CAC	GCG'	TGG	TGA	CCGI	CCC	CTC	CAG	CAGC	TI	YGGG	CAC	CC	AGAG	CT	CAT	CTG	CAA	CGTG
	TY	CCTY	CGC	ACC	ACTY	GGCA	CGG	GAC	GTC	CTCG	AA	CCC	GTG	GG	TCTC	3GAT	<b>I</b> GTA	GAC	GII	GCAC
188	S	S	V	V	T	V	P	S	s	s	L	G	T	Q	T	Y	I	С	N	V
1501	. A	ATC.	ACA	AGC	CCA	GCAA	CAC	CAA	\GG1	rcgac	AA	(GAA	AGT	TG	AGC	CCA	AATC	TIG	TGA	CAAA
	-TV	ראכי	ጥርሃፐ	TCG	CCT	CGTT	CTG	GTI	CCZ	AGCTG	T	CTI	<b>TCA</b>	AC	TCG	GT.	TTAG	AAC	ACT	G.L.I.I.
208	N	H	K	P	S	N	T	K	V	D	K	K	V	E	P	K	S	С	ט	K
1561					GCC															
					CGG			T												
228	T	H	T	C	P	P	0				_	5	3							

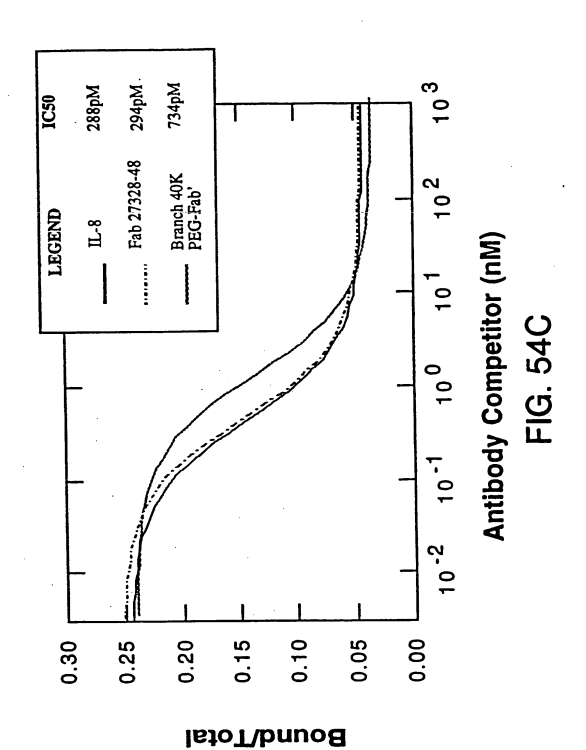
FIG. 53



**Bound/Total** 

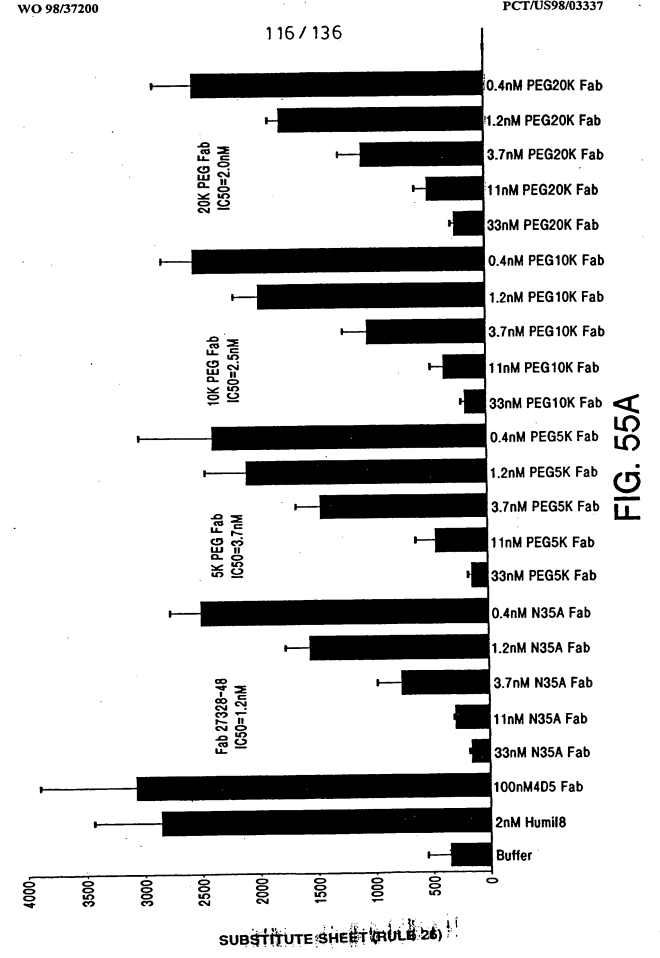


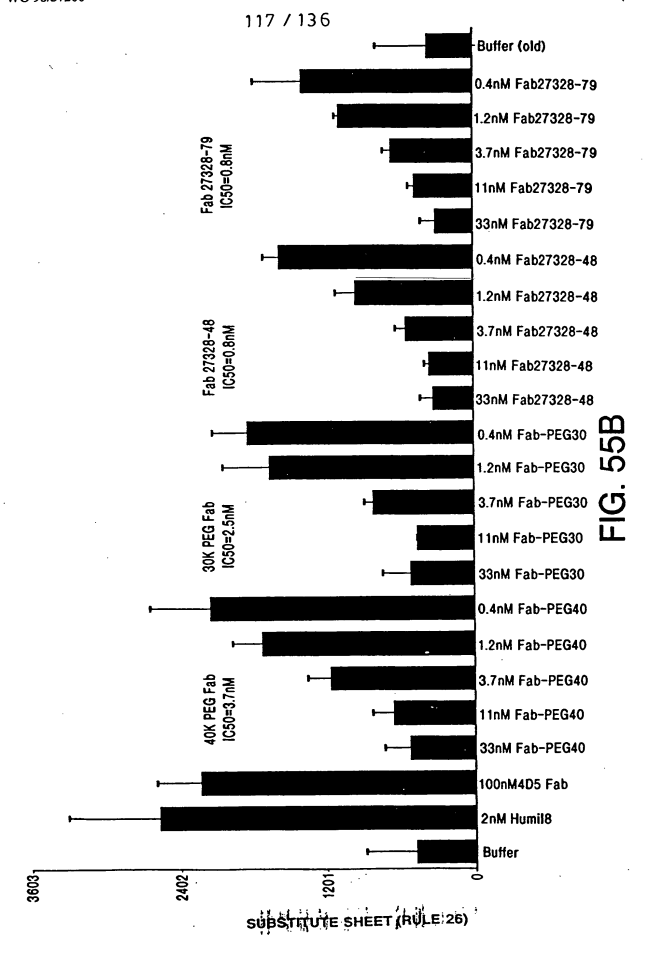
SUBSTITUTE SHEET (RULE 26)



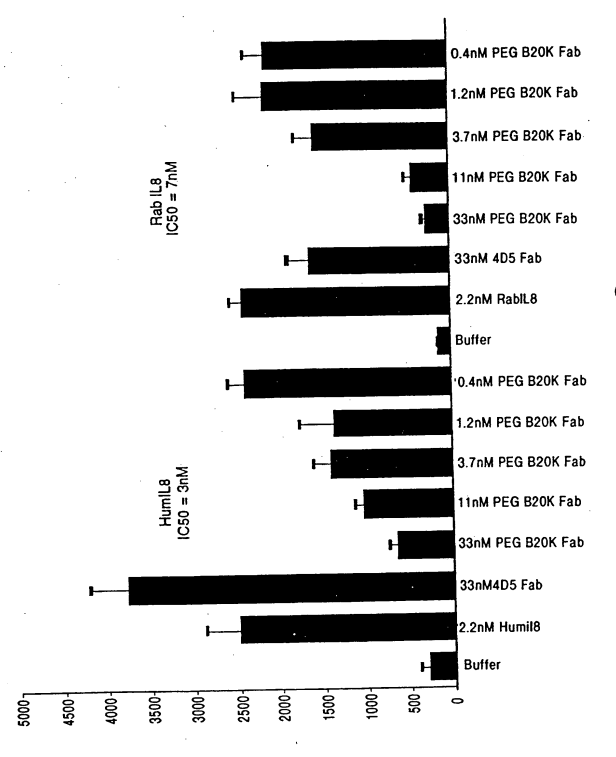
SUBSTITUTE SHEET (RULE 26)



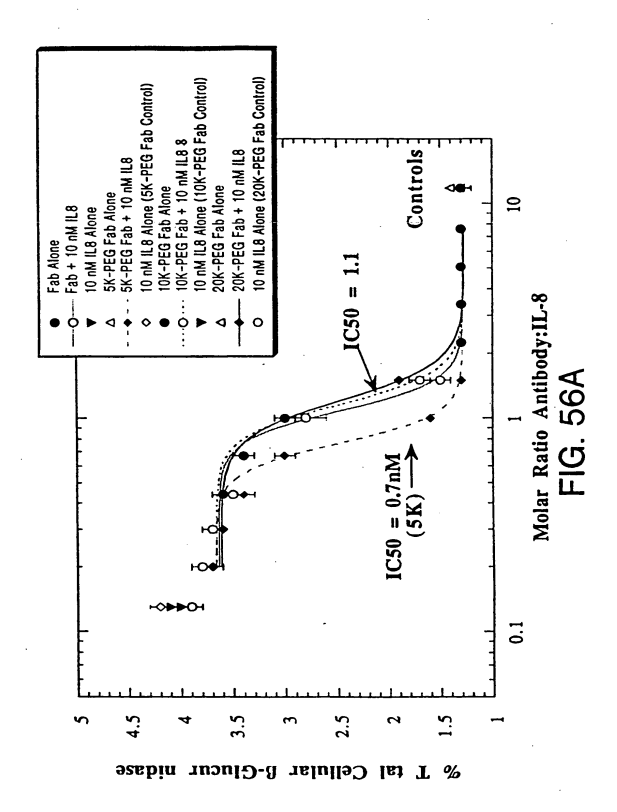


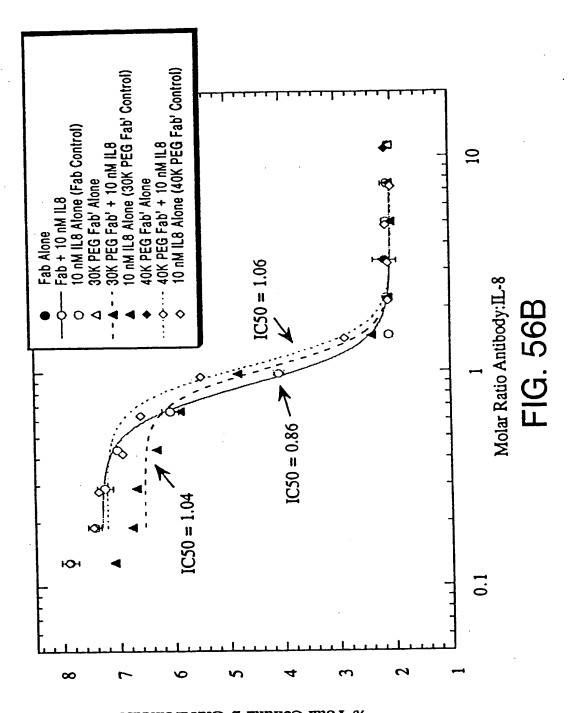


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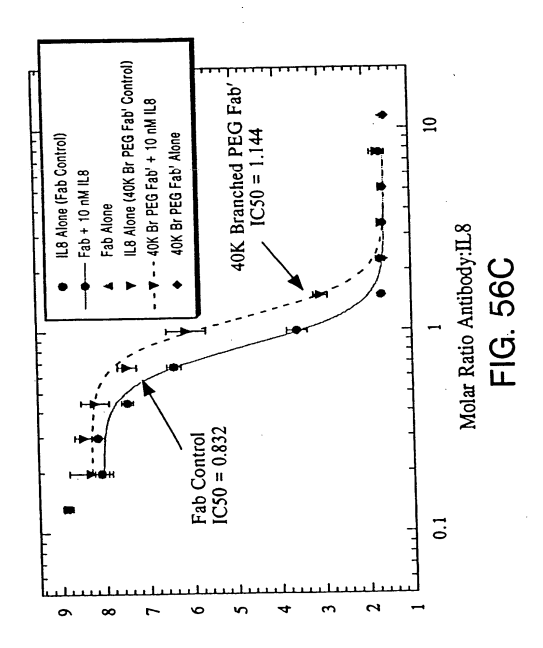


SUBSTITUTE SHEET (RULE 26)

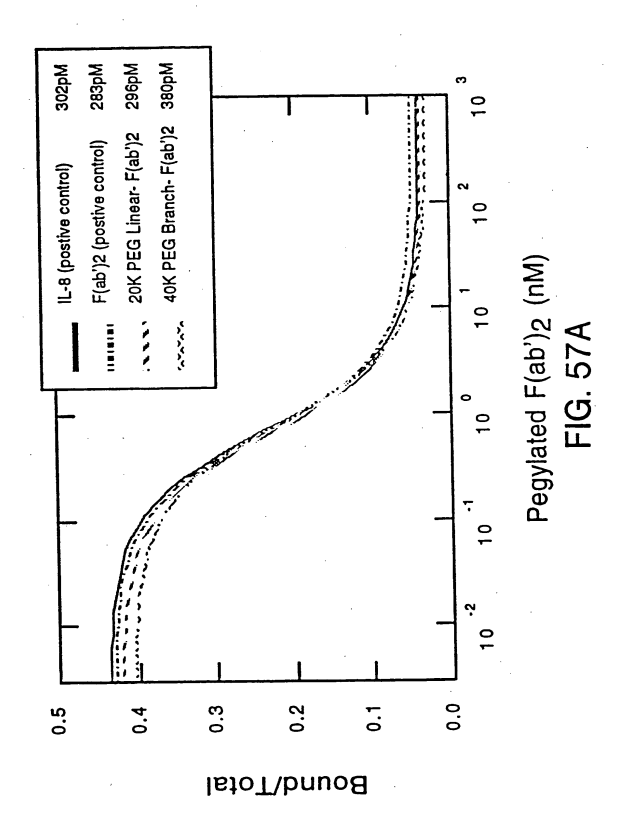




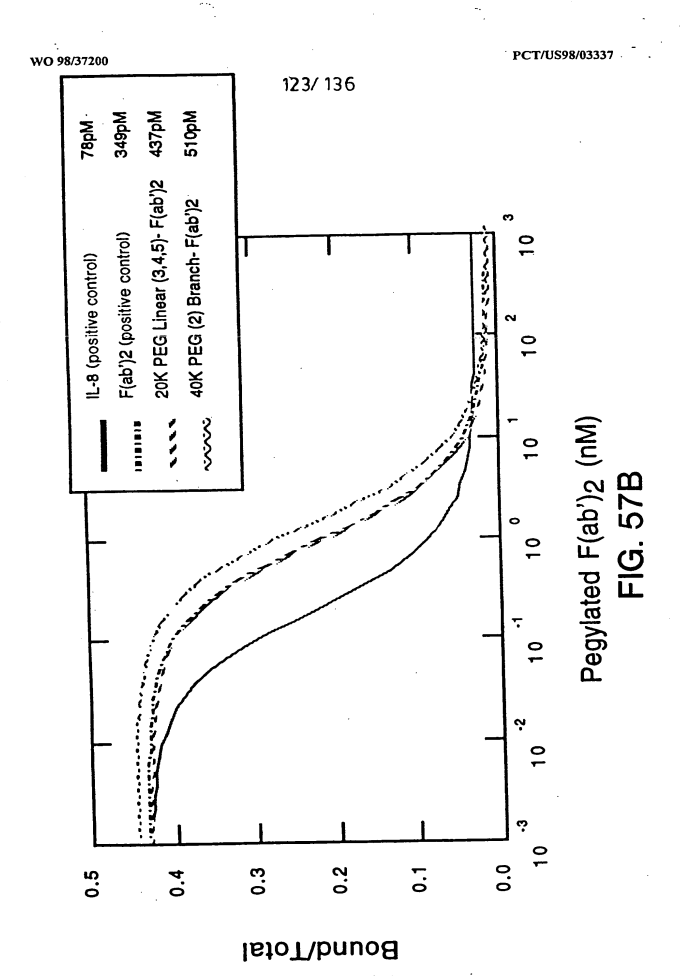
% Total Cellular B-Glucuronidase

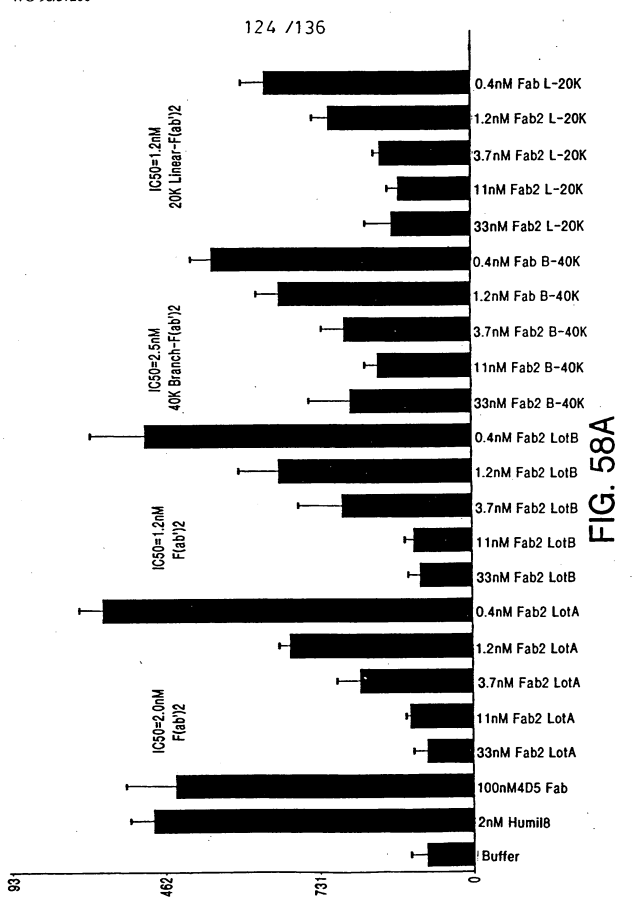


% Total Cellular B-Glucuronidase Activity



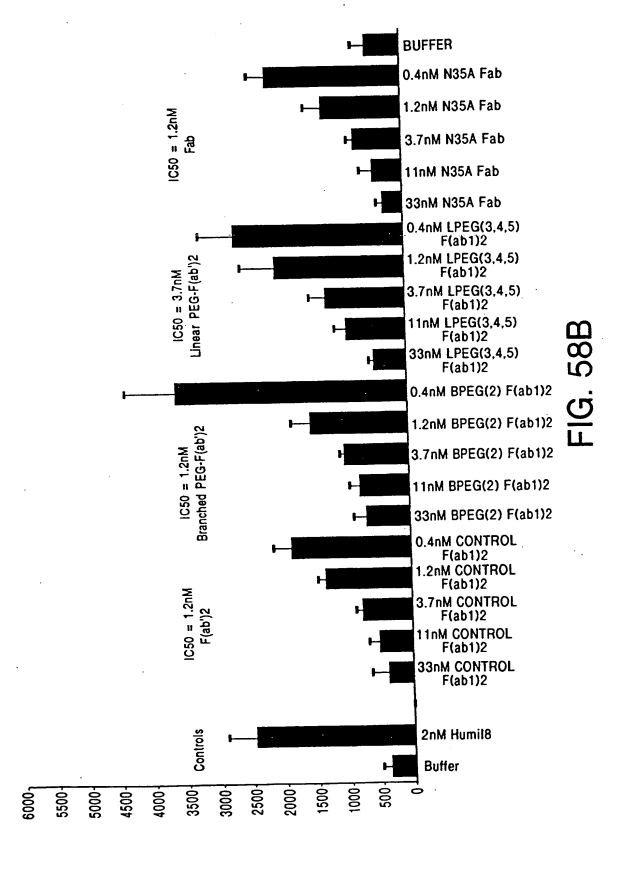
SUBSTITUTE SHEET (RULE 26)



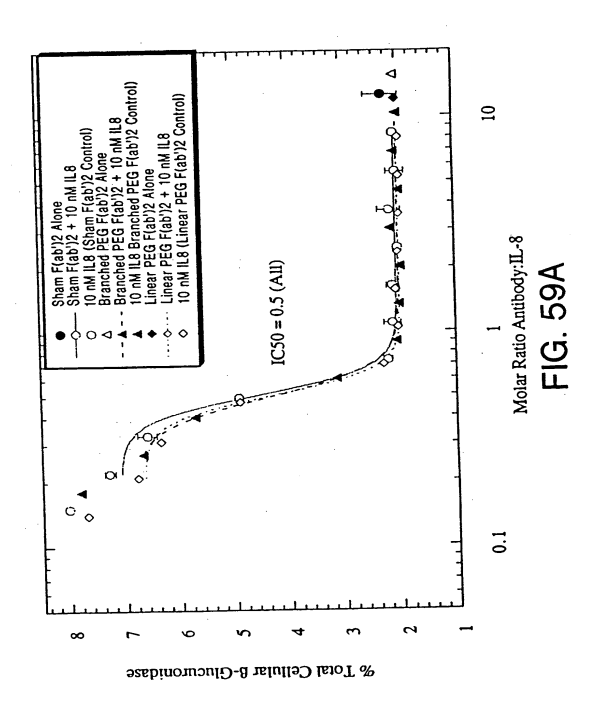


SUBSTITUTE SHEET (RULE 26)

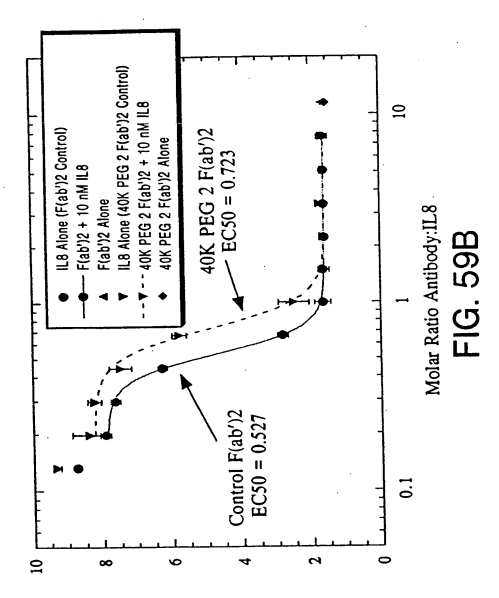
125/136



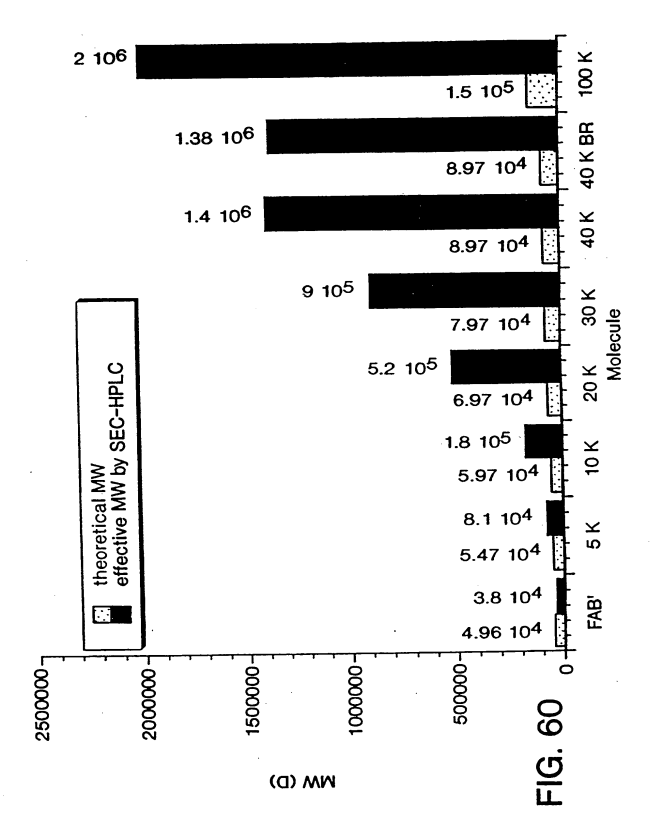
SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)



% Total Cellular B-Glucuronidase Activity



5K -10K -20K -30K -40K -40K branch -100K

Reduced



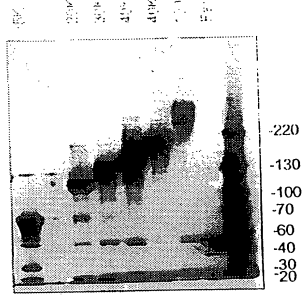
Fab-PEG-5000 -220 -130 -100

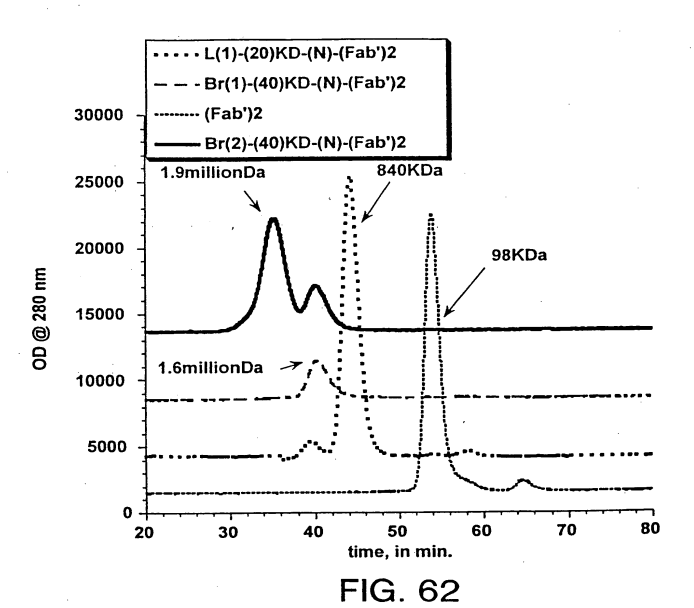
-70 -60 -40 -38 -20

FIG. 61A

Non-Reduced

FIG. 61B





SUBSTITUTE SHEET (RULE 26)

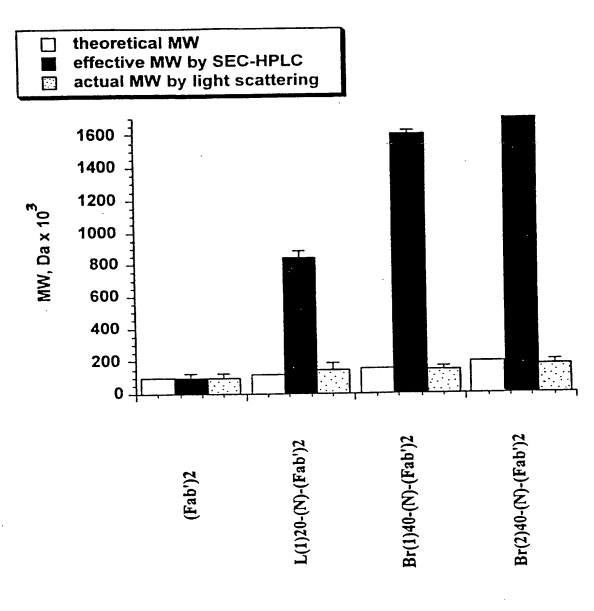


FIG. 63

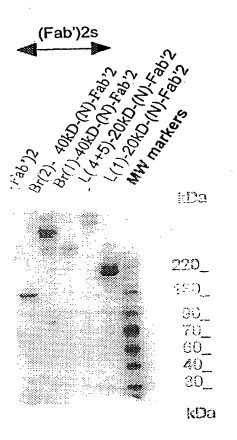
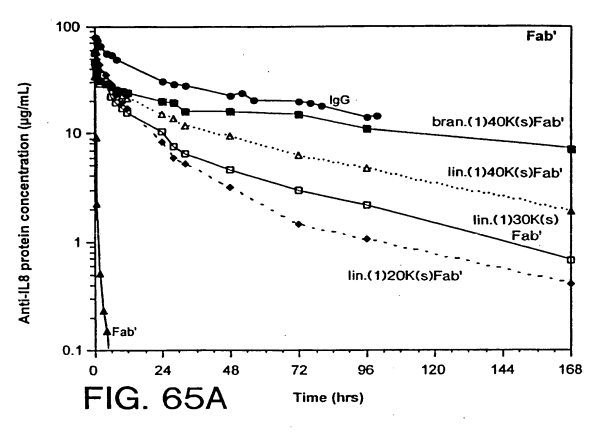
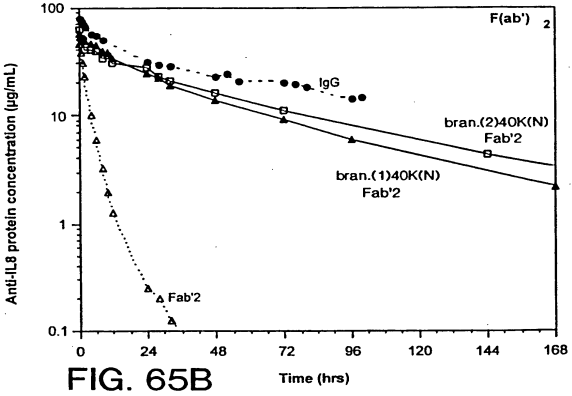


FIG. 64





SUBSTITUTE SHEET (RULE 26)

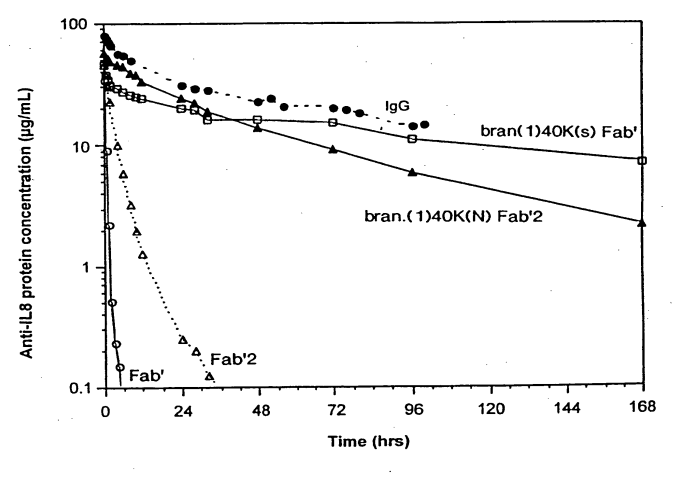
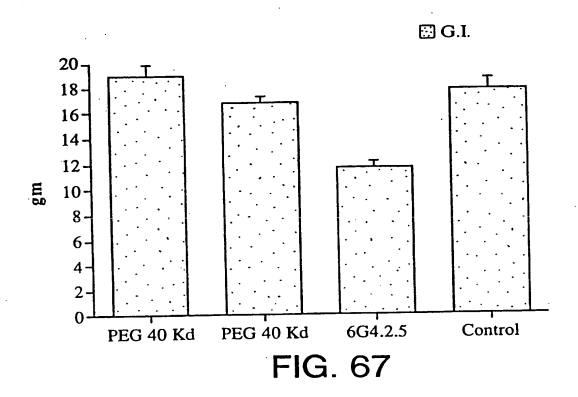
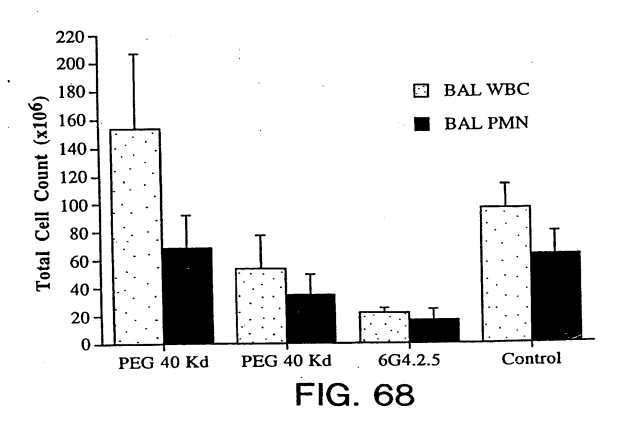
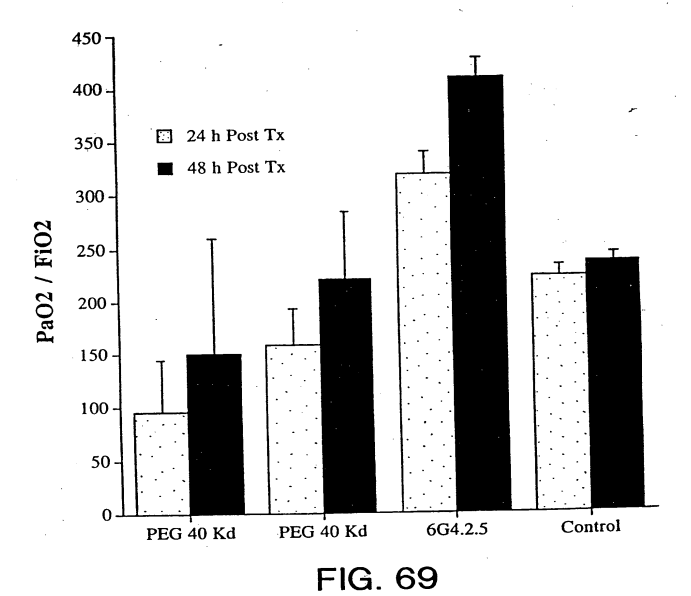


FIG. 66





SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)